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(54) Title: NUCLEIC ACIDS INCLUDING OPEN READING FRAMES ENCODING POLYPEPTIDES; "ORFX"

(57) Abstract

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The present invention provides open reading frames ORFX, encoding isolated polypeptides, as well as polynucleotides encoding ORFX and antibodies that immunospecifically bind to ORFX or any derivative, variant, mutant, or fragment of the ORFX polypeptides, polynucleotides or antibodies. The invention additionally provides methods in which the ORFX polypeptide, polynucleotide and antibody are used in detection and treatment of a broad range of pathological states, as well as to other uses.

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## NOVEL POLYNUCLEOTIDES AND POLYPEPTIDES ENCODED THEREBY

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## **BACKGROUND OF THE INVENTION**

The invention relates generally to nucleic acids and polypeptides encoded thereby, and methods of using these nucleic acids and polypeptides.

## SUMMARY OF THE INVENTION

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The invention is based in part on the discovery of nucleic acids that include open reading frames encoding novel polypeptides, and on the polypeptides encoded thereby. The nucleic acids and polypeptides are collectively referred to herein as "ORFX".

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Accordingly, in one aspect, the invention provides an isolated nucleic acid molecule (SEQ ID NO:2n-1, wherein n is an integer between 1-3161), that encodes novel polypeptide, or a fragment, homolog, analog or derivative thereof. The nucleic acid can include, e.g., a nucleic acid sequence encoding a polypeptide at least 85% identical to a polypeptide comprising the amino acid sequences of SEQ ID NO:2n, wherein n is an integer between 1-3161. The nucleic acid can be, e.g., a genomic DNA fragment, or a cDNA molecule.

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Also included in the invention is a vector containing one or more of the nucleic acids described herein, and a cell containing the vectors or nucleic acids described herein.

The invention is also directed to host cells transformed with a recombinant expression vector comprising any of the nucleic acid molecules described above.

In another aspect, the invention includes a pharmaceutical composition that includes an ORFX nucleic acid and a pharmaceutically acceptable carrier or diluent.

In a further aspect, the invention includes a substantially purified ORF polypeptide, e.g., any of the ORFX polypeptides encoded by an ORFX nucleic acid, and fragments, homologs, analogs, and derivatives thereof. The invention also includes a pharmaceutical composition that includes a ORFX polypeptide and a pharmaceutically acceptable carrier or diluent.

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In a still a further aspect, the invention provides an antibody that binds specifically to an ORFX polypeptide. The antibody can be, e.g., a monoclonal or polyclonal antibody, and fragments, homologs, analogs, and derivatives thereof. The invention also includes a pharmaceutical composition including ORFX antibody and a pharmaceutically acceptable carrier or diluent. The invention is also directed to isolated antibodies that bind to an epitope on a polypeptide encoded by any of the nucleic acid molecules described above.

The invention also includes kits comprising any of the pharmaceutical compositions described above.

The invention further provides a method for producing an ORFX polypeptide by providing a cell containing a ORFX nucleic acid, e.g., a vector that includes a ORFX nucleic acid, and culturing the cell under conditions sufficient to express the ORFX polypeptide encoded by the nucleic acid. The expressed ORFX polypeptide is then recovered from the cell. Preferably, the cell produces little or no endogenous ORFX polypeptide. The cell can be, e.g., a prokaryotic cell or eukaryotic cell.

The invention is also directed to methods of identifying an ORFX polypeptide or nucleic acids in a sample by contacting the sample with a compound that specifically binds to the polypeptide or nucleic acid, and detecting complex formation, if present.

The invention further provides methods of identifying a compound that modulates the activity of a ORFX polypeptide by contacting ORFX polypeptide with a compound and determining whether the ORFX polypeptide activity is modified.

The invention is also directed to compounds that modulate ORFX polypeptide activity identified by contacting a ORFX polypeptide with the compound and determining whether the compound modifies activity of the ORFX polypeptide, binds to the ORFX polypeptide, or binds to a nucleic acid molecule encoding a ORFX polypeptide.

In a another aspect, the invention provides a method of determining the presence of or predisposition of an ORFX-associated disorder in a subject. The method includes providing a sample from the subject and measuring the amount of ORFX polypeptide in the subject sample.

The amount of ORFX polypeptide in the subject sample is then compared to the amount of ORFX polypeptide in a control sample. An alteration in the amount of ORFX polypeptide in the subject protein sample relative to the amount of ORFX polypeptide in the control protein sample indicates the subject has a tissue proliferation-associated condition. A control sample is preferably taken from a matched individual, *i.e.*, an individual of similar age, sex, or other general condition but who is not suspected of having a tissue proliferation-associated condition. Alternatively, the control sample may be taken from the subject at a time when the subject is not suspected of having a tissue proliferation-associated disorder. In some embodiments, the ORFX is detected using a ORFX antibody.

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In a further aspect, the invention provides a method of determining the presence of or predisposition of an ORFX-associated disorder in a subject. The method includes providing a nucleic acid sample, e.g., RNA or DNA, or both, from the subject and measuring the amount of the ORFX nucleic acid in the subject nucleic acid sample. The amount of ORFX nucleic acid sample in the subject nucleic acid is then compared to the amount of an ORFX nucleic acid in a control sample. An alteration in the amount of ORFX nucleic acid in the sample relative to the amount of ORFX in the control sample indicates the subject has a tissue proliferation-associated disorder.

In a still further aspect, the invention provides method of treating or preventing or delaying a ORFX-associated disorder. The method includes administering to a subject in which such treatment or prevention or delay is desired a ORFX nucleic acid, a ORFX polypeptide, or an ORFX antibody in an amount sufficient to treat, prevent, or delay a tissue proliferation-associated disorder in the subject.

Unless otherwise defined, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. Although methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, suitable methods and materials are described below. All publications, patent applications, patents, and other references mentioned herein are incorporated by reference in their entirety. In the case of conflict, the present specification, including definitions, will control. In addition, the materials, methods, and examples are illustrative only and not intended to be limiting.

Other features and advantages of the invention will be apparent from the following detailed description and claims.

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# **DETAILED DESCRIPTION OF THE INVENTION**

The invention provides novel polypeptides and nucleotides encoded thereby. The polynucleotides and their encoded polypeptides can be grouped according to the functions played by their gene products. Such functions include, structural proteins, proteins from which associated with metabolic pathways fatty acid metabolism, glycolysis, intermediary metabolism, calcium metabolism, proteases, and amino acid metabolism, etc.

Included in the invention are 3161 novel nucleic acid sequences and their encoded polypeptides. The sequences are collectively referred to as "ORFX nucleic acids" or ORFX polynucleotides" and the corresponding encoded polypeptide is referred to as a "ORFX polypeptide" or ORFX protein". For example, an ORFX nucleic acid according to the invention is a nucleic acid including an ORF1 nucleic acid, and an ORF polypeptide according to the invention is a polypeptide that includes the amino acid sequence of an ORF1 polypeptide. Unless indicated otherwise, "ORFX" is meant to refer to any of the ORF1-3161 sequences disclosed herein.

Table 1 provides a summary of the ORFX nucleic acids and their encoded polypeptides are summarized in Table 1. Nucleic acid sequences and polypeptide sequences for ORFX nucleic acids according to the invention is provided in the section of the specification entitled "Disclosed Sequences of ORFX Nucleic Acid and Polypeptide Sequences."

Column 1 of Table 1, entitled "ORF #", denotes an ORF number assigned to a nucleic acid containing an open reading frame according to the invention.

Column 2 of Table 1, entitled "Internal Identification number (Nucleic Acid Sequence Identification Number, Polypeptide Sequence Identification Number), provides an internal identification number for the indicated ORF, along with sequence identification numbers (SEQ ID NOs.) corresponding to the indicated ORF. In general, for an ORFn according to the invention (wherein n is any integer from 1 to 3161), a nucleic acid corresponding to the ORF is SEQ ID NO:2n-1, and an amino acid sequence encoded by the ORF is SEQ ID NO:2n. For example, a nucleic acid sequence corresponding to an ORF1 nucleic acid is SEQ ID NO:1, and a polypeptide sequence corresponding to an ORF1 polypeptide is SEQ ID NO:2. Similarly, a

nucleic acid sequence corresponding to an ORF4 nucleic acid is SEQ ID NO:7, and a polypeptide sequence corresponding to an ORF4 polypeptide is SEQ ID NO:8; a nucleic acid sequence corresponding to an ORF198 nucleic acid sequence is SEQ ID NO:395, and a polypeptide sequence corresponding to an ORF198 polypeptide is SEQ ID NO:396. Nucleic acid sequences and polypeptide sequences for ORFX nucleic acids according to the invention are provided in the section of the specification entitled "Disclosed Sequences of ORFX Nucleic Acid and Polypeptide Sequences."

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Column 2 of Table 1, entitled "Protein Similarity", lists previously described proteins that are related to polypeptides encoded by the ORFs. Genbank identifiers for the previously described proteins are provided. These can be retrieved from <a href="http://www.ncbi.nlm.nih.gov/">http://www.ncbi.nlm.nih.gov/</a>.

To determine similarity to previously described proteins, polypeptides encoded by ORFX DNA sequences were tested using the Framesearch Algorithm against a nonredundant version of the GenPept Database from NCBI/Genbank. DNA sequences that had a score of '90' or above (Framesearch algorithm score, Edelman et. al. GCG Genetics) to a known protein were selected. Open reading frames were extended beyond the region of the protein matched using standard DNA translation and codon tables. Novel proteins that lacked a protein match were translated against the standard genetic codons and proteins with an ORF at least 80 amino acids and containing a Methionine start are included in the Table.

Column 3 of Table 3, entitled "Protein Domains", lists previously described protein domains, designated by pfam entries, that are present in polypeptides encoded by the ORFs, Also included in column 3 are proteins in which these domains are present. The pfam entries can be retrieved from <a href="http://pfam.wustl.edu/">http://pfam.wustl.edu/</a>. DNA sequences were translated in all six frames and tested using the Hmmer Algorithm against the Pfam Database (References to the algorithm and Pfam database can be found at <a href="http://pfam.wustl.edu">http://pfam.wustl.edu</a>). Translated DNA sequences that matched a protein domain entry in the Pfam database AND had a score of 7.5' were selected.

Column 4 of Table 3, entitled "Protein Classification", lists the type of classification assigned for the protein, based on its homology. Examples of proteins in the classification include the following proteins:

## **Amylases**

Amylase is responsible for endohydrolysis of 1,4-alpha-glucosidic linkages in oligosaccharides and polysaccharides. Variations in amylase gene may be indicative of delayed maturation and of various amylase producing neoplasms and carcinomas.

## 5 Amyloid

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The serum amyloid A (SAA) proteins comprise a family of vertebrate proteins that associate predominantly with high density lipoproteins (HDL). The synthesis of certain members of the family is greatly increased in inflammation. Prolonged elevation of plasma SAA levels, as in chronic inflammation, 15 results in a pathological condition, called amyloidosis, which affects the liver, kidney and spleen and which is characterized by the highly insoluble accumulation of SAA in these tissues. Amyloid selectively inhibits insulin-stimulated glucose utilization and glycogen deposition in muscle, while not affecting adipocyte glucose metabolism. Deposition of fibrillar amyloid proteins intraneuronally, as neurofibrillary tangles, extracellularly, as plaques and in blood vessels, is characteristic of both Alzheimer's disease and aged Down's syndrome. Amyloid deposition is also associated with type II diabetes mellitus.

## Angiopoeitin

Members of the angiopoeitin/fibrinogen family have been shown to stimulate the generation of new blood vessels, inhibit the generation of new blood vessels, and perform several roles in blood clotting. This generation of new blood vessels, called angiogenesis, is also an essential step in tumor growth in order for the tumor to get the blood supply it needs to expand. Variation in these genes may be predictive of any form of heart disease, numerous blood clotting disorders, stroke, hypertension and predisposition to tumor formation and metastasis. In particular, these variants may be predictive of the response to various antihypertensive drugs and chemotherapeutic and anti-tumor agents.

## Apoptosis-related proteins

Active cell suicide (apoptosis) is induced by events such as growth factor withdrawal and toxins. It is controlled by regulators, which have either an inhibitory effect on programmed cell

death (anti-apoptotic) or block the protective effect of inhibitors (pro-apoptotic). Many viruses have found a way of countering defensive apoptosis by encoding their own anti-apoptosis genes preventing their target-cells from dying too soon. Variants of apoptosis related genes may be useful in formulation of anti-aging drugs.

# Cadherin, Cyclin, Polymerase, Oncogenes, Histones, Kinases

Members of the cell division/cell cycle pathways such as cyclins, many transcription factors and kinases, DNA polymerases, histones, helicases and other oncogenes play a critical role in carcinogenesis where the uncontrolled proliferation of cells leads to tumor formation and eventually metastasis. Variation in these genes may be predictive of predisposition to any form of cancer, from increased risk of tumor formation to increased rate of metastasis. In particular, these variants may be predictive of the response to various chemotherapeutic and anti-tumor agents.

## Colony-stimulating factor-related proteins

Granulocyte/macrophage colony-stimulating factors are cytokines that act in hematopoiesis by controlling the production, differentiation, and function of 2 related white cell populations of the blood, the granulocytes and the monocytes-macrophages.

## Complement-related proteins

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Complement proteins are immune associated cytotoxic agents, acting in a chain reaction to exterminate target cells to that were opsonized (primed) with antibodies, by forming a membrane attack complex (MAC). The mechanism of killing is by opening pores in the target cell membrane. Variations in 20 complement genes or their inhibitors are associated with many autoimmune disorders. Modified serum levels of complement products cause edemas of various tissues, lupus (SLE), vasculitis, glomerulonephritis, renal failure, hemolytic anemia, thrombocytopenia, and arthritis. They interfere with mechanisms of ADCC (antibody dependent cell cytotoxicity), severely impair immune competence and reduce phagocytic ability. Variants of complement genes may also be indicative of type I diabetes mellitus, meningitis neurological disorders such as nemaline myopathy, neonatal hypotonia, muscular disorders such as congenital myopathy and other diseases.

## Cytochrome

The respiratory chain is a key biochemical pathway which is essential to all aerobic cells. There are five different cytochromes involved in the chain. These are heme bound proteins which serve as electron carriers. Modifications in these genes may be predictive of ataxia areflexia, dementia and myopathic and neuropathic changes in muscles. Also, association with various types of solid tumors.

#### **Kinesins**

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Kinesins are tubulin molecular motors that function to transport organelles within cells and to move chromosomes along microtubules during cell division. Modifications of these genes may be indicative of neurological disorders such as Pick disease of the brain, tuberous sclerosis.

# Cytokines, Interferon, Interleukin

Members of the cytokine families are known for their potent ability to stimulate cell growth and division even at low concentrations. Cytokines such as erythropoietin are cell-specific in their growth stimulation; erythropoietin is useful for the stimulation of the proliferation of erythroblasts. Variants in cytokines may be predictive for a wide variety of diseases, including cancer predisposition.

# G-protein coupled receptors

G-protein coupled receptors (also called R7G) are an extensive group of hormones, neurotransmitters, odorants and light receptors which transduce extracellular signals by interaction with guanine nucleotide-binding (G) proteins. Alterations in genes coding for G-coupled proteins may be involved in and indicative of a vast number of physiological conditions. These include blood pressure regulation, renal dysfunctions, male infertility, dopamine associated cognitive, emotional, and endocrine functions, hypercalcemia, chondrodysplasia and osteoporosis, pseudohypoparathyroidism, growth retardation and dwarfism.

## Thioesterases

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Eukaryotic thiol proteases are a family of proteolytic enzymes which contain an active site cysteine. Catalysis proceeds through a thioester intermediate and is facilitated by a nearby histidine side chain; an asparagine completes the essential catalytic triad. Variants of thioester associated genes may be predictive of neuronal disorders and mental illnesses such as Ceroid Lipoffiscinosis, Neuronal 1, Infantile, Santavuori disease and more.

The key to the molecule type is as follows:

10	Abbrev:	Title:
	amylase	amylase protein
	amylaseinhib	amylase inhibitor
	amyloid	amyloid protein
15	apoptosis	apoptosis associated protein
	apoptosisinhib	apoptosis inhibitors
	apoptosisrecep	apoptosis receptors
	ATPase_associated	ATPase associated protein
	biotindep	biotin dependent enzyme/protein
20	cadherin	cadherin protein
	calcium_channel	calcium channel protein
	carboxylase	carboxylase protein
	cathepsin	cathepsin/carboxypeptidases
	cathepsininhib	cathepsin/carboxypeptidase inhibitor
25	chloride_channel	chloride channel protein
	collagen	collagen
	complement	complement protein
	complementrecept	complement receptor protein
	complementinhib	complement inhibitor
30	csf	colony stimulating factor
	csfrecept	colony stimulating factor receptor
	cyclin	cyclin protein
	cyto450	cytochrome p450 protein
	cytochrome	cytochrome related protein
35	deaminase	deaminase
	dehydrogenase	dehydrogenase
	desaturase	desaturase
	dna_rna_bind	DNA/RNA binding protein/factor
	dna rna inhib	DNA/RNA binding protein/factor inhibitor
40	dynein	dynein
	•	•

	elastase elastaseinhib	elastase elastase inhibitor
•	eph esterase	EPH family of tyrosine kinases esterase
5	esteraseinhib	esterase inhibitor
	fgf	fibroblast growth factor
	fgfreceptor	fibroblast growth factor receptor
	gaba	GABA receptor
	glucoamylase	glucoamylase
10	glucoronidase	glucoronidase
	glycoprotein	glycoprotein
	Guanylyl	guanylylate cyclase
	helicase	helicase
	histone	histone
15	HOM	homologous
	homeobox	homeobox protein
	hydrolase	hydrolase
	hydroxysteroid	hydroxysteroid associated protein
	hypoxanthine	hypoxanthine associated protein
20	immunoglob	immunoglobulin
	immunoglobrecept	immunoglobulin receptor
	interferon	interferon
	interleukin	interleukin
	interleukinrecept	interleukin receptor
25	isomerase	isomerase
	isomeraseinhibitor	isomerase inhibitor
	isomerasereceptor	isomerase receptor kinase
	kinase kinaseinhibitor	kinase inhibitor
20		kinase receptor
30	kinasereceptor kinesin	kinesin
	laminin	laminin associated protein
	lipase	lipase
	metallothionein	metallothionein
35	MHC	major histocompatability complex
33	misc channel	miscellaneous channel
	ngf	nerve growth factor
	nuci_recpt	nuclear receptor
	nuclease	nuclease
40	oncogene	oncogene associated protein
•	oxidase	oxidase
	oxygenase	oxygenase
	peptidase	peptidase
	peroxidase	peroxidase
45	phosphatase	phosphatase
	phosphataseinhib	phosphatase inhibitor
	-	

	phosphorylase PIR	phosphorylase PIR DATABASE (release 56, 29-OCT- 1998)
	polymerase	polymerase
5	potassium_channel	potassium channel protein
•	prostaglandin	prostaglandin
	protease	protease
	proteaseinhib	protease inhibitor
	reductase	reductase
10	ribosomalprot	ribosomal associated protein
	RTR	EMBLDATABASE translated entries not to be incorporated into SWISS-PROT (20-JUL-1998)
	SIM	similar
15	SPTR	EMBL DATABASE translated entries to be incorporated into SWISS-PROT (20-JUL-1998)
	struct	structural associated protein
	sulfotransferase	sulfotransferase
20	SWP	SWISS-PROT DATABASE (release 18- OCT-1998)
	SWPN	SWISS-PROT Update (release 11-NOV-98)
	synthase	synthase
	tgf	transforming growth factor
25	tgfreceptor	transforming growth factor receptor
	thioesterase	thioesterase
	thiolase	thiolase
	tm7	seven transmembrane domain G-protein
		coupled receptor
30	tnf	necrosis factor receptor
	traffic	tumor necrosis factor
	tnfreceptor	tumor trafficking associated protein
	TRN	EMBL DATABASE translated entries
		update (20-JUL-1998)
35	transcriptfactor	transcription factor
	transferase	transferase
	transport	transport protein
	tubulin	tubulin
40	ubiquitin	ubiquitin
40	unclassified	Protein not categorized into one of the aforementioned protein families
	water channel	water channel protein

Column 5 of Table 1, entitled, "Cells or Tissues in Which Gene is Expressed", denotes tissues, represented by five digit numbers, in which RNA homologous to the ORF nucleic acid sequences is present. Tissues or cells corresponding to the numbers are provided in Table 2.

ORFX nucleic acids, and their encoded polypeptides, according to the invention are useful in a variety of applications and contexts. For example, various ORFX nucleic acids and polypeptides according to the invention are useful, *inter alia*, as novel members of the protein families indicated in Table 1, and/or according to the presence of domains and sequence relatedness to previously described proteins as summarized in Table 1.

ORFX nucleic acids and polypeptides according to the invention can also be used to identify cell types listed in Table 1 for an indicated ORFX according to the invention.

Additional utilities for ORFX nucleic acids and polypeptides according to the invention are disclosed herein.

#### **ORFX Nucleic Acids**

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The novel nucleic acids of the invention include those that encode an ORFX or ORFX-like protein, or biologically active portions thereof. The nucleic acids include nucleic acids encoding polypeptides that include the amino acid sequence of one or more of SEQ ID NO:2n, wherein n = 1 to 3161. The encoded polypeptides can thus include, e.g., the amino acid sequences of SEQ ID NO: 2, 4, 6, 8, 10, ..., 6310, 6312, 6314, 6316, 6318, 6320, and/or 6322.

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In some embodiments, a nucleic acid encoding a polypeptide having the amino acid sequence of one or more of SEQ ID NO:2n (wherein n = 1 to 3161) includes the nucleic acid sequence of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161), or a fragment thereof. Additionally, the invention includes mutant or variant nucleic acids of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161), or a fragment thereof, any of whose bases may be changed from the disclosed sequence while still encoding a protein that maintains its ORFX -like activities and physiological functions. The invention further includes the complement of the nucleic acid sequence of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161), including fragments, derivatives,

analogs and homolog thereof. The invention additionally includes nucleic acids or nucleic acid fragments, or complements thereto, whose structures include chemical modifications.

Also included are nucleic acid fragments sufficient for use as hybridization probes to identify ORFX-encoding nucleic acids (e.g., ORFX mRNA) and fragments for use as polymerase chain reaction (PCR) primers for the amplification or mutation of ORFX nucleic acid molecules. As used herein, the term "nucleic acid molecule" is intended to include DNA molecules (e.g., cDNA or genomic DNA), RNA molecules (e.g., mRNA), analogs of the DNA or RNA generated using nucleotide analogs, and derivatives, fragments and homologs thereof. The nucleic acid molecule can be single-stranded or double-stranded, but preferably is double-stranded DNA.

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"Probes" refer to nucleic acid sequences of variable length, preferably between at least about 10 nucleotides (nt), 100 nt, or as many as about, e.g., 6,000 nt, depending on use. Probes are used in the detection of identical, similar, or complementary nucleic acid sequences. Longer length probes are usually obtained from a natural or recombinant source, are highly specific and much slower to hybridize than oligomers. Probes may be single- or double-stranded and designed to have specificity in PCR, membrane-based hybridization technologies, or ELISA-like technologies.

An "isolated" nucleic acid molecule is one that is separated from other nucleic acid molecules that are present in the natural source of the nucleic acid. Examples of isolated nucleic acid molecules include, but are not limited to, recombinant DNA molecules contained in a vector, recombinant DNA molecules maintained in a heterologous host cell, partially or substantially purified nucleic acid molecules, and synthetic DNA or RNA molecules. Preferably, an "isolated" nucleic acid is free of sequences which naturally flank the nucleic acid (*i.e.*, sequences located at the 5' and 3' ends of the nucleic acid) in the genomic DNA of the organism from which the nucleic acid is derived. For example, in various embodiments, the isolated ORFX nucleic acid molecule can contain less than about 50 kb, 25 kb, 5 kb, 4 kb, 3 kb, 2 kb, 1 kb, 0.5 kb or 0.1 kb of nucleotide sequences which naturally flank the nucleic acid molecule in genomic DNA of the cell from which the nucleic acid is derived. Moreover, an "isolated" nucleic acid molecule, such as a cDNA molecule, can be substantially free of other cellular material or culture medium when produced by recombinant techniques, or of chemical precursors or other chemicals when chemically synthesized.

A nucleic acid molecule of the present invention, e.g., a nucleic acid molecule having the nucleotide sequence of SEQ ID NO:2n-1 (wherein n=1 to 3161), or a complement of any of this nucleotide sequence, can be isolated using standard molecular biology techniques and the sequence information provided herein. Using all or a portion of the nucleic acid sequence of any of SEQ ID NO:2n-1 (wherein n=1 to 3161) as a hybridization probe, ORFX nucleic acid sequences can be isolated using standard hybridization and cloning techniques (e.g., as described in Sambrook et al., eds., Molecular Cloning: A Laboratory Manual  $2^{nd}$  Ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, 1989; and Ausubel, et al., eds., Current Protocols in Molecular Biology, John Wiley & Sons, New York, NY, 1993.)

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A nucleic acid of the invention can be amplified using cDNA, mRNA or alternatively, genomic DNA, as a template and appropriate oligonucleotide primers according to standard PCR amplification techniques. The nucleic acid so amplified can be cloned into an appropriate vector and characterized by DNA sequence analysis. Furthermore, oligonucleotides corresponding to ORFX nucleotide sequences can be prepared by standard synthetic techniques, e.g., using an automated DNA synthesizer.

As used herein, the term "oligonucleotide" refers to a series of linked nucleotide residues, which oligonucleotide has a sufficient number of nucleotide bases to be used in a PCR reaction. A short oligonucleotide sequence may be based on, or designed from, a genomic or cDNA sequence and is used to amplify, confirm, or reveal the presence of an identical, similar or complementary DNA or RNA in a particular cell or tissue. Oligonucleotides comprise portions of a nucleic acid sequence having about 10 nt, 50 nt, or 100 nt in length, preferably about 15 nt to 30 nt in length. In one embodiment, an oligonucleotide comprising a nucleic acid molecule less than 100 nt in length would further comprise at lease 6 contiguous nucleotides of any of SEQ ID NO:2n-1 (wherein n=1 to 3161), or a complement thereof. Oligonucleotides may be chemically synthesized and may be used as probes.

In another embodiment, an isolated nucleic acid molecule of the invention comprises a nucleic acid molecule that is a complement of the nucleotide sequence shown in any of SEQ ID NO:2n-1 (wherein n=1 to 3161). In another embodiment, an isolated nucleic acid molecule of the invention comprises a nucleic acid molecule that is a complement of the nucleotide sequence shown in any of SEQ ID NO:2n-1 (wherein n=1 to 3161), or a portion of this nucleotide sequence. A nucleic acid molecule that is complementary to the nucleotide sequence shown in

is one that is sufficiently complementary to the nucleotide sequence shown in of any of SEQ ID NO:2n-1 (wherein n=1 to 3161) that it can hydrogen bond with little or no mismatches to the nucleotide sequence shown in of any of SEQ ID NO:2n-1 (wherein n=1 to 3161),, thereby forming a stable duplex.

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As used herein, the term "complementary" refers to Watson-Crick or Hoogsteen base pairing between nucleotides units of a nucleic acid molecule, and the term "binding" means the physical or chemical interaction between two polypeptides or compounds or associated polypeptides or compounds or combinations thereof. Binding includes ionic, non-ionic, Von der Waals, hydrophobic interactions, etc. A physical interaction can be either direct or indirect. Indirect interactions may be through or due to the effects of another polypeptide or compound. Direct binding refers to interactions that do not take place through, or due to, the effect of another polypeptide or compound, but instead are without other substantial chemical intermediates.

Moreover, the nucleic acid molecule of the invention can comprise only a portion of the nucleic acid sequence of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161), e.g., a fragment that can be used as a probe or primer, or a fragment encoding a biologically active portion of ORFX. Fragments provided herein are defined as sequences of at least 6 (contiguous) nucleic acids or at least 4 (contiguous) amino acids, a length sufficient to allow for specific hybridization in the case of nucleic acids or for specific recognition of an epitope in the case of amino acids, respectively, and are at most some portion less than a full length sequence. Fragments may be derived from any contiguous portion of a nucleic acid or amino acid sequence of choice. Derivatives are nucleic acid sequences or amino acid sequences formed from the native compounds either directly or by modification or partial substitution. Analogs are nucleic acid sequences or amino acid sequences that have a structure similar to, but not identical to, the native compound but differs from it in respect to certain components or side chains. Analogs may be synthetic or from a different evolutionary origin and may have a similar or opposite metabolic activity compared to wild type.

Derivatives and analogs may be full length or other than full length, if the derivative or analog contains a modified nucleic acid or amino acid, as described below. Derivatives or analogs of the nucleic acids or proteins of the invention include, but are not limited to, molecules comprising regions that are substantially homologous to the nucleic acids or proteins of the

invention, in various embodiments, by at least about 70%, 80%, 85%, 90%, 95%, 98%, or even 99% identity (with a preferred identity of 80-99%) over a nucleic acid or amino acid sequence of identical size or when compared to an aligned sequence in which the alignment is done by a computer homology program known in the art, or whose encoding nucleic acid is capable of hybridizing to the complement of a sequence encoding the aforementioned proteins under stringent, moderately stringent, or low stringent conditions. See e.g. Ausubel, et al., CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, John Wiley & Sons, New York, NY, 1993, and below. An exemplary program is the Gap program (Wisconsin Sequence Analysis Package, Version 8 for UNIX, Genetics Computer Group, University Research Park, Madison, WI) using the default settings, which uses the algorithm of Smith and Waterman (Adv. Appl. Math., 1981, 2: 482-489, which is incorporated herein by reference in its entirety).

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A "homologous nucleic acid sequence" or "homologous amino acid sequence," or variations thereof, refer to sequences characterized by a homology at the nucleotide level or amino acid level as discussed above. Homologous nucleotide sequences encode those sequences coding for isoforms of ORFX polypeptide. Isoforms can be expressed in different tissues of the same organism as a result of, for example, alternative splicing of RNA. Alternatively, isoforms can be encoded by different genes. In the present invention, homologous nucleotide sequences include nucleotide sequences encoding for a ORFX polypeptide of species other than humans, including, but not limited to, mammals, and thus can include, e.g., mouse, rat, rabbit, dog, cat cow, horse, and other organisms. Homologous nucleotide sequences also include, but are not limited to, naturally occurring allelic variations and mutations of the nucleotide sequences set forth herein. A homologous nucleotide sequence does not, however, include the nucleotide sequence encoding human ORFX protein. Homologous nucleic acid sequences include those nucleic acid sequences that encode conservative amino acid substitutions (see below) in any of SEQ ID NO:2n (wherein n = 1 to 3161) as well as a polypeptide having ORFX activity. Biological activities of the ORFX proteins are described below. A homologous amino acid sequence does not encode the amino acid sequence of a human ORFX polypeptide.

The nucleotide sequence determined from the cloning of the human ORFX gene allows for the generation of probes and primers designed for use in identifying the cell types disclosed and/or cloning ORFX homologues in other cell types, e.g., from other tissues, as well as ORFX homologues from other mammals. The probe/primer typically comprises a substantially purified

oligonucleotide. The oligonucleotide typically comprises a region of nucleotide sequence that hybridizes under stringent conditions to at least about 12, 25, 50, 100, 150, 200, 250, 300, 350 or 400 or more consecutive sense strand nucleotide sequence of SEQ ID NO:2n-1 (wherein n = 1 to 3161); or an anti-sense strand nucleotide sequence of SEQ ID NO:2n-1 (wherein n = 1 to 3161); or of a naturally occurring mutant of SEQ ID NO:2n-1 (wherein n = 1 to 3161).

Probes based on the human ORFX nucleotide sequence can be used to detect transcripts or genomic sequences encoding the same or homologous proteins. In various embodiments, the probe further comprises a label group attached thereto, e.g., the label group can be a radioisotope, a fluorescent compound, an enzyme, or an enzyme co-factor. Such probes can be used as a part of a diagnostic test kit for identifying cells or tissue which misexpress a ORFX protein, such as by measuring a level of a ORFX-encoding nucleic acid in a sample of cells from a subject e.g., detecting ORFX mRNA levels or determining whether a genomic ORFX gene has been mutated or deleted.

"A polypeptide having a biologically active portion of ORFX" refers to polypeptides exhibiting activity similar, but not necessarily identical to, an activity of a polypeptide of the present invention, including mature forms, as measured in a particular biological assay, with or without dose dependency. A nucleic acid fragment encoding a "biologically active portion of ORFX" can be prepared by isolating a portion of SEQ ID NO:2n-1 (wherein n=1 to 3161), that encodes a polypeptide having a ORFX biological activity (biological activities of the ORFX proteins are summarized in Table 1), expressing the encoded portion of ORFX protein (e.g., by recombinant expression in vitro) and assessing the activity of the encoded portion of ORFX. For example, a nucleic acid fragment encoding a biologically active portion of ORFX can optionally include a domain as shown in Table 1, column 4.

## ORFX variants

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The invention further encompasses nucleic acid molecules that differ from the disclosed ORFX nucleotide sequences due to degeneracy of the genetic code. These nucleic acids thus encode the same ORFX protein as that encoded by the nucleotide sequence shown in SEQ ID NO:2n-1 (wherein n=1 to 3161). In another embodiment, an isolated nucleic acid molecule of the invention has a nucleotide sequence encoding a protein having an amino acid sequence shown in any of SEQ ID NO:2n (wherein n=1 to 3161).

In addition to the human ORFX nucleotide sequence shown in any of SEQ ID NO:2n-1 (wherein n=1 to 3161), it will be appreciated by those skilled in the art that DNA sequence polymorphisms that lead to changes in the amino acid sequences of ORFX may exist within a population (e.g., the human population). Such genetic polymorphism in the ORFX gene may exist among individuals within a population due to natural allelic variation. As used herein, the terms "gene" and "recombinant gene" refer to nucleic acid molecules comprising an open reading frame encoding a ORFX protein, preferably a mammalian ORFX protein. Such natural allelic variations can typically result in 1-5% variance in the nucleotide sequence of the ORFX gene. Any and all such nucleotide variations and resulting amino acid polymorphisms in ORFX that are the result of natural allelic variation and that do not alter the functional activity of ORFX are intended to be within the scope of the invention.

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Moreover, nucleic acid molecules encoding ORFX proteins from other species, and thus that have a nucleotide sequence that differs from the human sequence of any of SEQ ID NO:2n-1 (wherein n=1 to 3161), are intended to be within the scope of the invention. Nucleic acid molecules corresponding to natural allelic variants and homologues of the ORFX cDNAs of the invention can be isolated based on their homology to the human ORFX nucleic acids disclosed herein using the human cDNAs, or a portion thereof, as a hybridization probe according to standard hybridization techniques under stringent hybridization conditions.

In another embodiment, an isolated nucleic acid molecule of the invention is at least 6 nucleotides in length and hybridizes under stringent conditions to the nucleic acid molecule comprising the nucleotide sequence of any of SEQ ID NO:2*n*-1 (wherein n = 1 to 3161). In another embodiment, the nucleic acid is at least 10, 25, 50, 100, 250, 500 or 750 nucleotides in length. In another embodiment, an isolated nucleic acid molecule of the invention hybridizes to the coding region. As used herein, the term "hybridizes under stringent conditions" is intended to describe conditions for hybridization and washing under which nucleotide sequences at least 60% homologous to each other typically remain hybridized to each other.

Homologs (i.e., nucleic acids encoding ORFX proteins derived from species other than human) or other related sequences (e.g., paralogs) can be obtained by low, moderate or high stringency hybridization with all or a portion of the particular human sequence as a probe using methods well known in the art for nucleic acid hybridization and cloning.

As used herein, the phrase "stringent hybridization conditions" refers to conditions under which a probe, primer or oligonucleotide will hybridize to its target sequence, but to no other sequences. Stringent conditions are sequence-dependent and will be different in different circumstances. Longer sequences hybridize specifically at higher temperatures than shorter sequences. Generally, stringent conditions are selected to be about 5°C lower than the thermal melting point (Tm) for the specific sequence at a defined ionic strength and pH. The Tm is the temperature (under defined ionic strength, pH and nucleic acid concentration) at which 50% of the probes complementary to the target sequence hybridize to the target sequence at equilibrium. Since the target sequences are generally present at excess, at Tm, 50% of the probes are occupied at equilibrium. Typically, stringent conditions will be those in which the salt concentration is less than about 1.0 M sodium ion, typically about 0.01 to 1.0 M sodium ion (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes, primers or oligonucleotides (e.g., 10 nt to 50 nt) and at least about 60°C for longer probes, primers and oligonucleotides. Stringent conditions may also be achieved with the addition of destabilizing agents, such as formamide.

Stringent conditions are known to those skilled in the art and can be found in CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, John Wiley & Sons, N.Y. (1989), 6.3.1-6.3.6. Preferably, the conditions are such that sequences at least about 65%, 70%, 75%, 85%, 90%, 95%, 98%, or 99% homologous to each other typically remain hybridized to each other. A non-limiting example of stringent hybridization conditions is hybridization in a high salt buffer comprising 6X SSC, 50 mM Tris-HCl (pH 7.5), 1 mM EDTA, 0.02% PVP, 0.02% Ficoll, 0.02% BSA, and 500 mg/ml denatured salmon sperm DNA at 65°C. This hybridization is followed by one or more washes in 0.2X SSC, 0.01% BSA at 50°C. An isolated nucleic acid molecule of the invention that hybridizes under stringent conditions to the sequence of any of SEQ ID NO:2n-1 (wherein n = 1 to 3161) corresponds to a naturally occurring nucleic acid molecule. As used herein, a "naturally-occurring" nucleic acid molecule refers to an RNA or DNA molecule having a nucleotide sequence that occurs in nature (e.g., encodes a natural protein).

In a second embodiment, a nucleic acid sequence that is hybridizable to the nucleic acid molecule comprising the nucleotide sequence of any of SEQ ID NO:2n-1 (wherein n=1 to 3161), or fragments, analogs or derivatives thereof, under conditions of moderate stringency is provided. A non-limiting example of moderate stringency hybridization conditions are

hybridization in 6X SSC, 5X Denhardt's solution, 0.5% SDS and 100 mg/ml denatured salmon sperm DNA at 55°C, followed by one or more washes in 1X SSC, 0.1% SDS at 37°C. Other conditions of moderate stringency that may be used are well known in the art. See, e.g., Ausubel et al. (eds.), 1993, Current Protocols in Molecular Biology, John Wiley & Sons, NY, and Kriegler, 1990, Gene Transfer and Expression, A Laboratory Manual, Stockton Press, NY.

In a third embodiment, a nucleic acid that is hybridizable to the nucleic acid molecule comprising the nucleotide sequence of any of SEQ ID NO:2*n*-1 (wherein *n* = 1 to 3161), or fragments, analogs or derivatives thereof, under conditions of low stringency, is provided. A non-limiting example of low stringency hybridization conditions are hybridization in 35% formamide, 5X SSC, 50 mM Tris-HCl (pH 7.5), 5 mM EDTA, 0.02% PVP, 0.02% Ficoll, 0.2% BSA, 100 mg/ml denatured salmon sperm DNA, 10% (wt/vol) dextran sulfate at 40°C, followed by one or more washes in 2X SSC, 25 mM Tris-HCl (pH 7.4), 5 mM EDTA, and 0.1% SDS at 50°C. Other conditions of low stringency that may be used are well known in the art (*e.g.*, as employed for cross-species hybridizations). See, *e.g.*, Ausubel *et al.* (eds.), 1993, CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, John Wiley & Sons, NY, and Kriegler, 1990, GENE TRANSFER AND EXPRESSION, A LABORATORY MANUAL, Stockton Press, NY; Shilo and Weinberg, 1981, *Proc Natl Acad Sci USA* 78: 6789-6792.

#### Conservative mutations

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In addition to naturally-occurring allelic variants of the ORFX sequence that may exist in the population, the skilled artisan will further appreciate that changes can be introduced by mutation into the nucleotide sequence of any of SEQ ID NO:2n-1 (wherein n=1 to 3161), thereby leading to changes in the amino acid sequence of the encoded ORFX protein, without altering the functional ability of the ORFX protein. For example, nucleotide substitutions leading to amino acid substitutions at "non-essential" amino acid residues can be made in the sequence of any of SEQ ID NO:2n-1 (wherein n=1 to 3161). A "non-essential" amino acid residue is a residue that can be altered from the wild-type sequence of ORFX without altering the biological activity, whereas an "essential" amino acid residue is required for biological activity. For example, amino acid residues that are conserved among the ORFX proteins of the present invention, are predicted to be particularly unamenable to alteration.

Amino acid residues that are conserved among members of an ORFX family members are predicted to be less amenable to alteration. For example, an ORFX protein according to the present invention can contain at least one domain (e.g., as shown in Table 1) that is a typically conserved region in an ORFX family member. As such, these conserved domains are not likely to be amenable to mutation. Other amino acid residues, however, (e.g., those that are not conserved or only semi-conserved among members of the ORFX family) may not be as essential for activity and thus are more likely to be amenable to alteration.

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Another aspect of the invention pertains to nucleic acid molecules encoding ORFX proteins that contain changes in amino acid residues that are not essential for activity. Such ORFX proteins differ in amino acid sequence from any of any of SEQ ID NO:2n (wherein n = 1 to 3161), yet retain biological activity. In one embodiment, the isolated nucleic acid molecule comprises a nucleotide sequence encoding a protein, wherein the protein comprises an amino acid sequence at least about 75% homologous to the amino acid sequence of any of SEQ ID NO:2n (wherein n = 1 to 3161). Preferably, the protein encoded by the nucleic acid is at least about 80% homologous to any of SEQ ID NO:2n (wherein n = 1 to 3161), more preferably at least about 90%, 95%, 98%, and most preferably at least about 99% homologous to SEQ ID NO:2.

An isolated nucleic acid molecule encoding a ORFX protein homologous to the protein of any of SEQ ID NO:2n (wherein n = 1 to 3161) can be created by introducing one or more nucleotide substitutions, additions or deletions into the corresponding nucleotide sequence, *i.e.* SEQ ID NO:2n-1 for the corresponding n, such that one or more amino acid substitutions, additions or deletions are introduced into the encoded protein.

Mutations can be introduced into SEQ ID NO:2n-1 (wherein n = 1 to 3161) by standard techniques, such as site-directed mutagenesis and PCR-mediated mutagenesis. Preferably, conservative amino acid substitutions are made at one or more predicted non-essential amino acid residues. A "conservative amino acid substitution" is one in which the amino acid residue is replaced with an amino acid residue having a similar side chain. Families of amino acid residues having similar side chains have been defined in the art. These families include amino acids with basic side chains (e.g., lysine, arginine, histidine), acidic side chains (e.g., aspartic acid, glutamic acid), uncharged polar side chains (e.g., glycine, asparagine, glutamine, serine, threonine, tyrosine, cysteine), nonpolar side chains (e.g., alanine, valine, leucine, isoleucine, proline,

phenylalanine, methionine, tryptophan), beta-branched side chains (e.g., threonine, valine, isoleucine) and aromatic side chains (e.g., tyrosine, phenylalanine, tryptophan, histidine). Thus, a predicted nonessential amino acid residue in ORFX is replaced with another amino acid residue from the same side chain family. Alternatively, in another embodiment, mutations can be introduced randomly along all or part of a ORFX coding sequence, such as by saturation mutagenesis, and the resultant mutants can be screened for ORFX biological activity to identify mutants that retain activity. Following mutagenesis of SEQ ID NO:2n-1 (wherein n = 1 to 3161), the encoded protein can be expressed by any recombinant technology known in the art and the activity of the protein can be determined.

In one embodiment, a mutant ORFX protein can be assayed for (1) the ability to form protein:protein interactions with other ORFX proteins, other cell-surface proteins, or biologically active portions thereof, (2) complex formation between a mutant ORFX protein and a ORFX receptor; (3) the ability of a mutant ORFX protein to bind to an intracellular target protein or biologically active portion thereof; (e.g., avidin proteins); (4) the ability to bind BRA protein; or (5) the ability to specifically bind an anti-ORFX protein antibody.

## Antisense

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Another aspect of the invention pertains to isolated antisense nucleic acid molecules that are hybridizable to or complementary to the nucleic acid molecule comprising the nucleotide sequence of SEQ ID NO:2n-1 (wherein n = 1 to 3161), or fragments, analogs or derivatives thereof. An "antisense" nucleic acid comprises a nucleotide sequence that is complementary to a "sense" nucleic acid encoding a protein, e.g., complementary to the coding strand of a double-stranded cDNA molecule or complementary to an mRNA sequence. In specific aspects, antisense nucleic acid molecules are provided that comprise a sequence complementary to at least about 10, 25, 50, 100, 250 or 500 nucleotides or an entire ORFX coding strand, or to only a portion thereof. Nucleic acid molecules encoding fragments, homologs, derivatives and analogs of a ORFX protein of any of SEQ ID NO:2n (wherein n = 1 to 3161) or antisense nucleic acids complementary to a ORFX nucleic acid sequence of SEQ ID NO:2n-1 (wherein n = 1 to 3161) are additionally provided.

In one embodiment, an antisense nucleic acid molecule is antisense to a "coding region" of the coding strand of a nucleotide sequence encoding ORFX. The term "coding region" refers to the region of the nucleotide sequence comprising codons which are translated into amino acid

residues (e.g., the protein coding region of a human ORFX that corresponds to any of SEQ ID NO:2n (wherein n = 1 to 3161)). In another embodiment, the antisense nucleic acid molecule is antisense to a "noncoding region" of the coding strand of a nucleotide sequence encoding ORFX. The term "noncoding region" refers to 5' and 3' sequences which flank the coding region that are not translated into amino acids (i.e., also referred to as 5' and 3' untranslated regions).

Given the coding strand sequences encoding ORFX disclosed herein (e.g., SEQ ID NO:2n-1 (wherein n = 1 to 3161)), antisense nucleic acids of the invention can be designed according to the rules of Watson and Crick or Hoogsteen base pairing. The antisense nucleic acid molecule can be complementary to the entire coding region of ORFX mRNA, but more preferably is an oligonucleotide that is antisense to only a portion of the coding or noncoding region of ORFX mRNA. For example, the antisense oligonucleotide can be complementary to the region surrounding the translation start site of ORFX mRNA. An antisense oligonucleotide can be, for example, about 5, 10, 15, 20, 25, 30, 35, 40, 45 or 50 nucleotides in length. An antisense nucleic acid of the invention can be constructed using chemical synthesis or enzymatic ligation reactions using procedures known in the art. For example, an antisense nucleic acid (e.g., an antisense oligonucleotide) can be chemically synthesized using naturally occurring nucleotides or variously modified nucleotides designed to increase the biological stability of the molecules or to increase the physical stability of the duplex formed between the antisense and sense nucleic acids, e.g., phosphorothioate derivatives and acridine substituted nucleotides can be used.

Examples of modified nucleotides that can be used to generate the antisense nucleic acid include: 5-fluorouracil, 5-bromouracil, 5-chlorouracil, 5-iodouracil, hypoxanthine, xanthine, 4-acetylcytosine, 5-(carboxyhydroxylmethyl) uracil, 5-carboxymethylaminomethyl-2-thiouridine, 5-carboxymethylaminomethyluracil, dihydrouracil, beta-D-galactosylqueosine, inosine, N6-isopentenyladenine, 1-methylguanine, 1-methylinosine, 2,2-dimethylguanine, 2-methyladenine, 2-methylguanine, 3-methylcytosine, 5-methylcytosine, N6-adenine, 7-methylguanine, 5-methyluracil, 5-methoxyaminomethyl-2-thiouracil, beta-D-mannosylqueosine, 5'-methoxycarboxymethyluracil, 5-methoxyuracil, 2-methylthio-N6-isopentenyladenine, uracil-5-oxyacetic acid (v), wybutoxosine, pseudouracil, queosine, 2-thiocytosine, 5-methyl-2-thiouracil, 2-thiouracil, 4-thiouracil, 5-methyluracil, uracil-5-oxyacetic acid (v), 5-methyl-2-thiouracil,

3-(3-amino-3-N-2-carboxypropyl) uracil, (acp3)w, and 2,6-diaminopurine. Alternatively, the antisense nucleic acid can be produced biologically using an expression vector into which a nucleic acid has been subcloned in an antisense orientation (*i.e.*, RNA transcribed from the inserted nucleic acid will be of an antisense orientation to a target nucleic acid of interest, described further in the following subsection).

The antisense nucleic acid molecules of the invention are typically administered to a subject or generated in situ such that they hybridize with or bind to cellular mRNA and/or genomic DNA encoding a ORFX protein to thereby inhibit expression of the protein, e.g., by inhibiting transcription and/or translation. The hybridization can be by conventional nucleotide complementarity to form a stable duplex, or, for example, in the case of an antisense nucleic acid molecule that binds to DNA duplexes, through specific interactions in the major groove of the double helix. An example of a route of administration of antisense nucleic acid molecules of the invention includes direct injection at a tissue site. Alternatively, antisense nucleic acid molecules can be modified to target selected cells and then administered systemically. For example, for systemic administration, antisense molecules can be modified such that they specifically bind to receptors or antigens expressed on a selected cell surface, e.g., by linking the antisense nucleic acid molecules to peptides or antibodies that bind to cell surface receptors or antigens. The antisense nucleic acid molecules can also be delivered to cells using the vectors described herein. To achieve sufficient intracellular concentrations of antisense molecules, vector constructs in which the antisense nucleic acid molecule is placed under the control of a strong pol II or pol III promoter are preferred.

In yet another embodiment, the antisense nucleic acid molecule of the invention is an α-anomeric nucleic acid molecule. An α-anomeric nucleic acid molecule forms specific double-stranded hybrids with complementary RNA in which, contrary to the usual β-units, the strands run parallel to each other (Gaultier et al. (1987) Nucleic Acids Res 15: 6625-6641). The antisense nucleic acid molecule can also comprise a 2'-o-methylribonucleotide (Inoue et al. (1987) Nucleic Acids Res 15: 6131-6148) or a chimeric RNA -DNA analogue (Inoue et al. (1987) FEBS Lett 215: 327-330).

## Ribozymes and PNA moieties

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Such modifications include, by way of nonlimiting example, modified bases, and nucleic acids whose sugar phosphate backbones are modified or derivatized. These modifications are

carried out at least in part to enhance the chemical stability of the modified nucleic acid, such that they may be used, for example, as antisense binding nucleic acids in therapeutic applications in a subject.

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In still another embodiment, an antisense nucleic acid of the invention is a ribozyme. Ribozymes are catalytic RNA molecules with ribonuclease activity that are capable of cleaving a single-stranded nucleic acid, such as an mRNA, to which they have a complementary region. Thus, ribozymes (e.g., hammerhead ribozymes (described in Haselhoff and Gerlach (1988) Nature 334:585-591)) can be used to catalytically cleave ORFX mRNA transcripts to thereby inhibit translation of ORFX mRNA. A ribozyme having specificity for a ORFX-encoding nucleic acid can be designed based upon the nucleotide sequence of a ORFX DNA disclosed herein (i.e., SEQ ID NO:2n-1 (wherein n = 1 to 3161)). For example, a derivative of a Tetrahymena L-19 IVS RNA can be constructed in which the nucleotide sequence of the active site is complementary to the nucleotide sequence to be cleaved in a ORFX-encoding mRNA. See, e.g., Cech et al. U.S. Pat. No. 4,987,071; and Cech et al. U.S. Pat. No. 5,116,742. Alternatively, ORFX mRNA can be used to select a catalytic RNA having a specific ribonuclease activity from a pool of RNA molecules. See, e.g., Bartel et al., (1993) Science 261:1411-1418.

Alternatively, ORFX gene expression can be inhibited by targeting nucleotide sequences complementary to the regulatory region of the ORFX (e.g., the ORFX promoter and/or enhancers) to form triple helical structures that prevent transcription of the ORFX gene in target cells. See generally, Helene. (1991) Anticancer Drug Des. 6: 569-84; Helene. et al. (1992) Ann. N.Y. Acad. Sci. 660:27-36; and Maher (1992) Bioassays 14: 807-15.

In various embodiments, the nucleic acids of ORFX can be modified at the base moiety, sugar moiety or phosphate backbone to improve, e.g., the stability, hybridization, or solubility of the molecule. For example, the deoxyribose phosphate backbone of the nucleic acids can be modified to generate peptide nucleic acids (see Hyrup et al. (1996) Bioorg Med Chem 4: 5-23). As used herein, the terms "peptide nucleic acids" or "PNAs" refer to nucleic acid mimics, e.g., DNA mimics, in which the deoxyribose phosphate backbone is replaced by a pseudopeptide backbone and only the four natural nucleobases are retained. The neutral backbone of PNAs has been shown to allow for specific hybridization to DNA and RNA under conditions of low ionic strength. The synthesis of PNA oligomers can be performed using standard solid phase peptide

synthesis protocols as described in Hyrup et al. (1996) above; Perry-O'Keefe et al. (1996) PNAS 93: 14670-675.

PNAs of ORFX can be used in therapeutic and diagnostic applications. For example, PNAs can be used as antisense or antigene agents for sequence-specific modulation of gene expression by, e.g., inducing transcription or translation arrest or inhibiting replication. PNAs of ORFX can also be used, e.g., in the analysis of single base pair mutations in a gene by, e.g., PNA directed PCR clamping; as artificial restriction enzymes when used in combination with other enzymes, e.g., S1 nucleases (Hyrup B. (1996) above); or as probes or primers for DNA sequence and hybridization (Hyrup et al. (1996), above; Perry-O'Keefe (1996), above).

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In another embodiment, PNAs of ORFX can be modified, e.g., to enhance their stability or cellular uptake, by attaching lipophilic or other helper groups to PNA, by the formation of PNA-DNA chimeras, or by the use of liposomes or other techniques of drug delivery known in the art. For example, PNA-DNA chimeras of ORFX can be generated that may combine the advantageous properties of PNA and DNA. Such chimeras allow DNA recognition enzymes, e.g., RNase H and DNA polymerases, to interact with the DNA portion while the PNA portion would provide high binding affinity and specificity. PNA-DNA chimeras can be linked using linkers of appropriate lengths selected in terms of base stacking, number of bonds between the nucleobases, and orientation (Hyrup (1996) above). The synthesis of PNA-DNA chimeras can be performed as described in Hyrup (1996) above and Finn et al. (1996) Nucl Acids Res 24: 3357-63. For example, a DNA chain can be synthesized on a solid support using standard phosphoramidite coupling chemistry, and modified nucleoside analogs, e.g., 5'-(4-methoxytrityl)amino-5'-deoxy-thymidine phosphoramidite, can be used between the PNA and the 5' end of DNA (Mag et al. (1989) Nucl Acid Res 17: 5973-88). PNA monomers are then coupled in a stepwise manner to produce a chimeric molecule with a 5' PNA segment and a 3' DNA segment (Finn et al. (1996) above). Alternatively, chimeric molecules can be synthesized with a 5' DNA segment and a 3' PNA segment. See, Petersen et al. (1975) Bioorg Med Chem Lett 5: 1119-11124.

In other embodiments, the oligonucleotide may include other appended groups such as peptides (e.g., for targeting host cell receptors in vivo), or agents facilitating transport across the cell membrane (see, e.g., Letsinger et al., 1989, Proc. Natl. Acad. Sci. U.S.A. 86:6553-6556; Lemaitre et al., 1987, Proc. Natl. Acad. Sci. 84:648-652; PCT Publication No. W088/09810) or

the blood-brain barrier (see, e.g., PCT Publication No. W089/10134). In addition, oligonucleotides can be modified with hybridization triggered cleavage agents (See, e.g., Krol et al., 1988, BioTechniques 6:958-976) or intercalating agents. (See, e.g., Zon, 1988, Pharm. Res. 5: 539-549). To this end, the oligonucleotide may be conjugated to another molecule, e.g., a peptide, a hybridization triggered cross-linking agent, a transport agent, a hybridization-triggered cleavage agent, etc.

## **ORFX** polypeptides

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The novel protein of the invention includes the ORFX-like protein whose sequence is provided in any of SEQ ID NO:2n (wherein n = 1 to 3161). The invention also includes a mutant or variant protein any of whose residues may be changed from the corresponding residue shown in FIG. 1 while still encoding a protein that maintains its ORFX-like activities and physiological functions, or a functional fragment thereof. For example, the invention includes the polypeptides encoded by the variant ORFX nucleic acids described above. In the mutant or variant protein, up to 20% or more of the residues may be so changed.

In general, an ORFX -like variant that preserves ORFX-like function includes any variant in which residues at a particular position in the sequence have been substituted by other amino acids, and further include the possibility of inserting an additional residue or residues between two residues of the parent protein as well as the possibility of deleting one or more residues from the parent sequence. Any amino acid substitution, insertion, or deletion is encompassed by the invention. In favorable circumstances, the substitution is a conservative substitution as defined above. Furthermore, without limiting the scope of the invention, positions of any of SEQ ID NO:2n (wherein n=1 to 3161) may be substitute such that a mutant or variant protein may include one or more substitutions

The invention also includes isolated ORFX proteins, and biologically active portions thereof, or derivatives, fragments, analogs or homologs thereof. Also provided are polypeptide fragments suitable for use as immunogens to raise anti-ORFX antibodies. In one embodiment, native ORFX proteins can be isolated from cells or tissue sources by an appropriate purification scheme using standard protein purification techniques. In another embodiment, ORFX proteins are produced by recombinant DNA techniques. Alternative to recombinant expression, a ORFX

protein or polypeptide can be synthesized chemically using standard peptide synthesis techniques.

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An "isolated" or "purified" protein or biologically active portion thereof is substantially free of cellular material or other contaminating proteins from the cell or tissue source from which the ORFX protein is derived, or substantially free from chemical precursors or other chemicals when chemically synthesized. The language "substantially free of cellular material" includes preparations of ORFX protein in which the protein is separated from cellular components of the cells from which it is isolated or recombinantly produced. In one embodiment, the language "substantially free of cellular material" includes preparations of ORFX protein having less than about 30% (by dry weight) of non-ORFX protein (also referred to herein as a "contaminating protein"), more preferably less than about 20% of non-ORFX protein, still more preferably less than about 10% of non-ORFX protein, and most preferably less than about 5% non-ORFX protein. When the ORFX protein or biologically active portion thereof is recombinantly produced, it is also preferably substantially free of culture medium, *i.e.*, culture medium represents less than about 20%, more preferably less than about 10%, and most preferably less than about 5% of the volume of the protein preparation.

The language "substantially free of chemical precursors or other chemicals" includes preparations of ORFX protein in which the protein is separated from chemical precursors or other chemicals that are involved in the synthesis of the protein. In one embodiment, the language "substantially free of chemical precursors or other chemicals" includes preparations of ORFX protein having less than about 30% (by dry weight) of chemical precursors or non-ORFX chemicals, more preferably less than about 20% chemical precursors or non-ORFX chemicals, still more preferably less than about 10% chemical precursors or non-ORFX chemicals, and most preferably less than about 5% chemical precursors or non-ORFX chemicals.

Biologically active portions of a ORFX protein include peptides comprising amino acid sequences sufficiently homologous to or derived from the amino acid sequence of the ORFX protein, e.g., the amino acid sequence shown in SEQ ID NO:2 that include fewer amino acids than the full length ORFX proteins, and exhibit at least one activity of a ORFX protein. Typically, biologically active portions comprise a domain or motif with at least one activity of the ORFX protein. A biologically active portion of a ORFX protein can be a polypeptide which is, for example, 10, 25, 50, 100 or more amino acids in length.

A biologically active portion of a ORFX protein of the present invention may contain at least one of the above-identified domains conserved between the FGF family of proteins. Moreover, other biologically active portions, in which other regions of the protein are deleted, can be prepared by recombinant techniques and evaluated for one or more of the functional activities of a native ORFX protein.

In an embodiment, the ORFX protein has an amino acid sequence shown in any of SEQ ID NO:2n (wherein n = 1 to 3161). In other embodiments, the ORFX protein is substantially homologous to any of SEQ ID NO:2n (wherein n = 1 to 3161) and retains the functional activity of the protein of any of SEQ ID NO:2n (wherein n = 1 to 3161), yet differs in amino acid sequence due to natural allelic variation or mutagenesis, as described in detail below. Accordingly, in another embodiment, the ORFX protein is a protein that comprises an amino acid sequence at least about 45% homologous, and more preferably about 55, 65, 70, 75, 80, 85, 90, 95, 98 or even 99% homologous to the amino acid sequence of any of SEQ ID NO:2n (wherein n = 1 to 3161) and retains the functional activity of the ORFX proteins of the corresponding polypeptide having the sequence of SEQ ID NO:2n (wherein n = 1 to 3161).

## Determining homology between two or more sequences

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To determine the percent homology of two amino acid sequences or of two nucleic acids, the sequences are aligned for optimal comparison purposes (e.g., gaps can be introduced in either of the sequences being compared for optimal alignment between the sequences). The amino acid residues or nucleotides at corresponding amino acid positions or nucleotide positions are then compared. When a position in the first sequence is occupied by the same amino acid residue or nucleotide as the corresponding position in the second sequence, then the molecules are homologous at that position (i.e., as used herein amino acid or nucleic acid "homology" is equivalent to amino acid or nucleic acid "identity").

The nucleic acid sequence homology may be determined as the degree of identity between two sequences. The homology may be determined using computer programs known in the art, such as GAP software provided in the GCG program package. See, *Needleman and Wunsch* 1970 *J Mol Biol* 48: 443-453. Using GCG GAP software with the following settings for nucleic acid sequence comparison: GAP creation penalty of 5.0 and GAP extension penalty of 0.3, the coding region of the analogous nucleic acid sequences referred to above exhibits a

degree of identity preferably of at least 70%, 75%, 80%, 85%, 90%, 95%, 98%, or 99%, with the CDS (encoding) part of the DNA sequence shown in SEQ ID NO:2n-1 (wherein n = 1 to 3161).

The term "sequence identity" refers to the degree to which two polynucleotide or polypeptide sequences are identical on a residue-by-residue basis over a particular region of comparison. The term "percentage of sequence identity" is calculated by comparing two optimally aligned sequences over that region of comparison, determining the number of positions at which the identical nucleic acid base (e.g., A, T, C, G, U, or I, in the case of nucleic acids) occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the region of comparison (i.e., the window size), and multiplying the result by 100 to yield the percentage of sequence identity. The term "substantial identity" as used herein denotes a characteristic of a polynucleotide sequence, wherein the polynucleotide comprises a sequence that has at least 80 percent sequence identity, preferably at least 85 percent identity and often 90 to 95 percent sequence identity, more usually at least 99 percent sequence identity as compared to a reference sequence over a comparison region. The term "percentage of positive residues" is calculated by comparing two optimally aligned sequences over that region of comparison, determining the number of positions at which the identical and conservative amino acid substitutions, as defined above, occur in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the region of comparison (i.e., the window size), and multiplying the result by 100 to yield the percentage of positive residues.

## Chimeric and fusion proteins

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The invention also provides ORFX chimeric or fusion proteins. As used herein, a ORFX "chimeric protein" or "fusion protein" includes a ORFX polypeptide operatively linked to a non-ORFX polypeptide. A "ORFX polypeptide" refers to a polypeptide having an amino acid sequence corresponding to ORFX, whereas a "non-ORFX polypeptide" refers to a polypeptide having an amino acid sequence corresponding to a protein that is not substantially homologous to the ORFX protein, e.g., a protein that is different from the ORFX protein and that is derived from the same or a different organism. Within a ORFX fusion protein the ORFX polypeptide can correspond to all or a portion of a ORFX protein. In one embodiment, a ORFX fusion protein comprises at least one biologically active portion of a ORFX protein. In another embodiment, a ORFX fusion protein comprises at least two biologically active portions of a

ORFX protein. Within the fusion protein, the term "operatively linked" is intended to indicate that the ORFX polypeptide and the non-ORFX polypeptide are fused in-frame to each other. The non-ORFX polypeptide can be fused to the N-terminus or C-terminus of the ORFX polypeptide.

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For example, in one embodiment a ORFX fusion protein comprises a ORFX polypeptide operably linked to the extracellular domain of a second protein. Such fusion proteins can be further utilized in screening assays for compounds that modulate ORFX activity (such assays are described in detail below).

In another embodiment, the fusion protein is a GST-ORFX fusion protein in which the ORFX sequences are fused to the C-terminus of the GST (i.e., glutathione S-transferase) sequences. Such fusion proteins can facilitate the purification of recombinant ORFX.

In yet another embodiment, the fusion protein is a ORFX protein containing a heterologous signal sequence at its N-terminus. For example, the native ORFX signal sequence can be removed and replaced with a signal sequence from another protein. In certain host cells (e.g., mammalian host cells), expression and/or secretion of ORFX can be increased through use of a heterologous signal sequence.

In another embodiment, the fusion protein is a ORFX-immunoglobulin fusion protein in which the ORFX sequences comprising one or more domains are fused to sequences derived from a member of the immunoglobulin protein family. The ORFX-immunoglobulin fusion proteins of the invention can be incorporated into pharmaceutical compositions and administered to a subject to inhibit an interaction between a ORFX ligand and a ORFX protein on the surface of a cell, to thereby suppress ORFX-mediated signal transduction *in vivo*. In one nonlimiting example, a contemplated ORFX ligand of the invention is an ORFX receptor. The ORFX-immunoglobulin fusion proteins can be used to modulate the bioavailability of a ORFX cognate ligand. Inhibition of the ORFX ligand/ORFX interaction may be useful therapeutically for both the treatment of proliferative and differentiative disorders, as well as modulating (e.g., promoting or inhibiting) cell survival. Moreover, the ORFX-immunoglobulin fusion proteins of the invention can be used as immunogens to produce anti-ORFX antibodies in a subject, to purify ORFX ligands, and in screening assays to identify molecules that inhibit the interaction of ORFX with a ORFX ligand.

A ORFX chimeric or fusion protein of the invention can be produced by standard recombinant DNA techniques. For example, DNA fragments coding for the different polypeptide sequences are ligated together in-frame in accordance with conventional techniques, e.g., by employing blunt-ended or stagger-ended termini for ligation, restriction enzyme digestion to provide for appropriate termini, filling-in of cohesive ends as appropriate, alkaline phosphatase treatment to avoid undesirable joining, and enzymatic ligation. In another embodiment, the fusion gene can be synthesized by conventional techniques including automated DNA synthesizers. Alternatively, PCR amplification of gene fragments can be carried out using anchor primers that give rise to complementary overhangs between two consecutive gene fragments that can subsequently be annealed and reamplified to generate a chimeric gene sequence (see, for example, Ausubel et al. (eds.) CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, John Wiley & Sons, 1992). Moreover, many expression vectors are commercially available that already encode a fusion moiety (e.g., a GST polypeptide). A ORFX-encoding nucleic acid can be cloned into such an expression vector such that the fusion moiety is linked in-frame to the ORFX protein.

## **ORFX** agonists and antagonists

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The present invention also pertains to variants of the ORFX proteins that function as either ORFX agonists (mimetics) or as ORFX antagonists. Variants of the ORFX protein can be generated by mutagenesis, e.g., discrete point mutation or truncation of the ORFX protein. An agonist of the ORFX protein can retain substantially the same, or a subset of, the biological activities of the naturally occurring form of the ORFX protein. An antagonist of the ORFX protein can inhibit one or more of the activities of the naturally occurring form of the ORFX protein by, for example, competitively binding to a downstream or upstream member of a cellular signaling cascade which includes the ORFX protein. Thus, specific biological effects can be elicited by treatment with a variant of limited function. In one embodiment, treatment of a subject with a variant having a subset of the biological activities of the naturally occurring form of the protein has fewer side effects in a subject relative to treatment with the naturally occurring form of the ORFX proteins.

Variants of the ORFX protein that function as either ORFX agonists (mimetics) or as ORFX antagonists can be identified by screening combinatorial libraries of mutants, e.g., truncation mutants, of the ORFX protein for ORFX protein agonist or antagonist activity. In one

embodiment, a variegated library of ORFX variants is generated by combinatorial mutagenesis at the nucleic acid level and is encoded by a variegated gene library. A variegated library of ORFX variants can be produced by, for example, enzymatically ligating a mixture of synthetic oligonucleotides into gene sequences such that a degenerate set of potential ORFX sequences is expressible as individual polypeptides, or alternatively, as a set of larger fusion proteins (e.g., for phage display) containing the set of ORFX sequences therein. There are a variety of methods which can be used to produce libraries of potential ORFX variants from a degenerate oligonucleotide sequence. Chemical synthesis of a degenerate gene sequence can be performed in an automatic DNA synthesizer, and the synthetic gene then ligated into an appropriate expression vector. Use of a degenerate set of genes allows for the provision, in one mixture, of all of the sequences encoding the desired set of potential ORFX sequences. Methods for synthesizing degenerate oligonucleotides are known in the art (see, e.g., Narang (1983) Tetrahedron 39:3; Itakura et al. (1984) Annu Rev Biochem 53:323; Itakura et al. (1984) Science 198:1056; Ike et al. (1983) Nucl Acid Res 11:477.

## Polypeptide libraries

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In addition, libraries of fragments of the ORFX protein coding sequence can be used to generate a variegated population of ORFX fragments for screening and subsequent selection of variants of a ORFX protein. In one embodiment, a library of coding sequence fragments can be generated by treating a double stranded PCR fragment of a ORFX coding sequence with a nuclease under conditions wherein nicking occurs only about once per molecule, denaturing the double stranded DNA, renaturing the DNA to form double stranded DNA that can include sense/antisense pairs from different nicked products, removing single stranded portions from reformed duplexes by treatment with S1 nuclease, and ligating the resulting fragment library into an expression vector. By this method, an expression library can be derived which encodes N-terminal and internal fragments of various sizes of the ORFX protein.

Several techniques are known in the art for screening gene products of combinatorial libraries made by point mutations or truncation, and for screening cDNA libraries for gene products having a selected property. Such techniques are adaptable for rapid screening of the gene libraries generated by the combinatorial mutagenesis of ORFX proteins. The most widely used techniques, which are amenable to high throughput analysis, for screening large gene libraries typically include cloning the gene library into replicable expression vectors,

transforming appropriate cells with the resulting library of vectors, and expressing the combinatorial genes under conditions in which detection of a desired activity facilitates isolation of the vector encoding the gene whose product was detected. Recrusive ensemble mutagenesis (REM), a new technique that enhances the frequency of functional mutants in the libraries, can be used in combination with the screening assays to identify ORFX variants (Arkin and Yourvan (1992) PNAS 89:7811-7815; Delgrave *et al.* (1993) Protein Engineering 6:327-331).

## **Anti-ORFX Antibodies**

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The invention further encompasses antibodies and antibody fragments, such as  $F_{ab}$  or  $(F_{ab})_2$  that bind immunospecifically to any of the proteins of the invention.

An isolated ORFX protein, or a portion or fragment thereof, can be used as an immunogen to generate antibodies that bind ORFX using standard techniques for polyclonal and monoclonal antibody preparation. Full-length ORFX protein can be used. Alternatively, the invention provides antigenic peptide fragments of ORFX for use as immunogens. The antigenic peptide of ORFX comprises at least 4 amino acid residues of the amino acid sequence shown in any of SEQ ID NO:2n (wherein n = 1 to 3161). The antigenic peptide encompasses an epitope of ORFX such that an antibody raised against the peptide forms a specific immune complex with ORFX. The antigenic peptide may comprise at least 6 aa residues, at least 8 aa residues, at least 10 aa residues, at least 15 aa residues, at least 20 aa residues, or at least 30 aa residues. In one embodiment of the invention, the antigenic peptide comprises a polypeptide comprising at least 6 contiguous amino acids of any of SEQ ID NO:2n (wherein n = 1 to 3161).

In an embodiment of the invention, epitopes encompassed by the antigenic peptide are regions of ORFX that are located on the surface of the protein, e.g., hydrophilic regions. As a means for targeting antibody production, hydropathy plots showing regions of hydrophilicity and hydrophobicity may be generated by any method well known in the art, including, for example, the Kyte Doolittle or the Hopp Woods methods, either with or without Fourier transformation. See, e.g., Hopp and Woods, 1981, Proc. Nat. Acad. Sci. USA 78: 3824-3828; Kyte and Doolittle 1982, J. Mol. Biol. 157: 105-142, each incorporated herein by reference in their entirety.

As disclosed herein, an ORFX protein sequence of any of SEQ ID NO:2n (wherein n = 1 to 3161), or derivatives, fragments, analogs or homologs thereof, may be utilized as immunogens in the generation of antibodies that immunospecifically-bind these protein components. The term "antibody" as used herein refers to immunoglobulin molecules and

immunologically active portions of immunoglobulin molecules, i.e., molecules that contain an antigen binding site that specifically binds (immunoreacts with) an antigen, such as ORFX. Such antibodies include, but are not limited to, polyclonal, monoclonal, chimeric, single chain,  $F_{ab}$  and  $F_{(ab')2}$  fragments, and an  $F_{ab}$  expression library. In a specific embodiment, antibodies to human ORFX proteins are disclosed. Various procedures known within the art may be used for the production of polyclonal or monoclonal antibodies to a ORFX protein sequence of any of SEQ ID NO:2n (wherein n = 1 to 3161) or derivative, fragment, analog or homolog thereof. Some of these proteins are discussed below.

For the production of polyclonal antibodies, various suitable host animals (e.g., rabbit, goat, mouse or other mammal) may be immunized by injection with the native protein, or a synthetic variant thereof, or a derivative of the foregoing. An appropriate immunogenic preparation can contain, for example, recombinantly expressed ORFX protein or a chemically synthesized ORFX polypeptide. The preparation can further include an adjuvant. Various adjuvants used to increase the immunological response include, but are not limited to, Freund's (complete and incomplete), mineral gels (e.g., aluminum hydroxide), surface active substances (e.g., lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, dinitrophenol, etc.), human adjuvants such as Bacille Calmette-Guerin and Corynebacterium parvum, or similar immunostimulatory agents. If desired, the antibody molecules directed against ORFX can be isolated from the mammal (e.g., from the blood) and further purified by well known techniques, such as protein A chromatography to obtain the IgG fraction.

The term "monoclonal antibody" or "monoclonal antibody composition", as used herein, refers to a population of antibody molecules that contain only one species of an antigen binding site capable of immunoreacting with a particular epitope of ORFX. A monoclonal antibody composition thus typically displays a single binding affinity for a particular ORFX protein with which it immunoreacts. For preparation of monoclonal antibodies directed towards a particular ORFX protein, or derivatives, fragments, analogs or homologs thereof, any technique that provides for the production of antibody molecules by continuous cell line culture may be utilized. Such techniques include, but are not limited to, the hybridoma technique (see Kohler & Milstein, 1975 *Nature* 256: 495-497); the trioma technique; the human B-cell hybridoma technique (see Kozbor, *et al.*, 1983 *Immunol Today* 4: 72) and the EBV hybridoma technique to produce human monoclonal antibodies (see Cole, *et al.*, 1985 In: MONOCLONAL ANTIBODIES AND

CANCER THERAPY, Alan R. Liss, Inc., pp. 77-96). Human monoclonal antibodies may be utilized in the practice of the present invention and may be produced by using human hybridomas (see Cote, et al., 1983. Proc Natl Acad Sci USA 80: 2026-2030) or by transforming human B-cells with Epstein Barr Virus in vitro (see Cole, et al., 1985 In: MONOCLONAL ANTIBODIES AND CANCER THERAPY, Alan R. Liss, Inc., pp. 77-96). Each of the above citations are incorporated herein by reference in their entirety

According to the invention, techniques can be adapted for the production of single-chain antibodies specific to a ORFX protein (see e.g., U.S. Patent No. 4,946,778). In addition, methods can be adapted for the construction of  $F_{ab}$  expression libraries (see e.g., Huse, et al., 1989 Science 246: 1275-1281) to allow rapid and effective identification of monoclonal  $F_{ab}$  fragments with the desired specificity for a ORFX protein or derivatives, fragments, analogs or homologs thereof. Non-human antibodies can be "humanized" by techniques well known in the art. See e.g., U.S. Patent No. 5,225,539. Each of the above citations are incorporated herein by reference. Antibody fragments that contain the idiotypes to a ORFX protein may be produced by techniques known in the art including, but not limited to: (i) an  $F_{(ab)/2}$  fragment produced by pepsin digestion of an antibody molecule; (ii) an  $F_{ab}$  fragment generated by reducing the disulfide bridges of an  $F_{(ab)/2}$  fragment; (iii) an  $F_{ab}$  fragment generated by the treatment of the antibody molecule with papain and a reducing agent and (iv)  $F_{v}$  fragments.

Additionally, recombinant anti-ORFX antibodies, such as chimeric and humanized monoclonal antibodies, comprising both human and non-human portions, which can be made using standard recombinant DNA techniques, are within the scope of the invention. Such chimeric and humanized monoclonal antibodies can be produced by recombinant DNA techniques known in the art, for example using methods described in PCT International Application No. PCT/US86/02269; European Patent Application No. 184,187; European Patent Application No. 171,496; European Patent Application No. 173,494; PCT International Publication No. WO 86/01533; U.S. Pat. No. 4,816,567; European Patent Application No. 125,023; Better et al.(1988) Science 240:1041-1043; Liu et al. (1987) PNAS 84:3439-3443; Liu et al. (1987) J Immunol. 139:3521-3526; Sun et al. (1987) PNAS 84:214-218; Nishimura et al. (1987) Cancer Res 47:999-1005; Wood et al. (1985) Nature 314:446-449; Shaw et al. (1988), J. Natl Cancer Inst 80:1553-1559); Morrison(1985) Science 229:1202-1207; Oi et al. (1986) BioTechniques 4:214; U.S. Pat. No. 5,225,539; Jones et al. (1986) Nature 321:552-525;

Verhoeyan et al. (1988) Science 239:1534; and Beidler et al. (1988) J Immunol 141:4053-4060. Each of the above citations are incorporated herein by reference.

In one embodiment, methods for the screening of antibodies that possess the desired specificity include, but are not limited to, enzyme-linked immunosorbent assay (ELISA) and other immunologically-mediated techniques known within the art. In a specific embodiment, selection of antibodies that are specific to a particular domain of a ORFX protein is facilitated by generation of hybridomas that bind to the fragment of a ORFX protein possessing such a domain. Antibodies that are specific for one or more domains within a ORFX protein, e.g., the domain spanning the first fifty amino-terminal residues specific to ORFX when compared to FGF-9, or derivatives, fragments, analogs or homologs thereof, are also provided herein.

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Anti-ORFX antibodies may be used in methods known within the art relating to the localization and/or quantitation of a ORFX protein (e.g., for use in measuring levels of the ORFX protein within appropriate physiological samples, for use in diagnostic methods, for use in imaging the protein, and the like). In a given embodiment, antibodies for ORFX proteins, or derivatives, fragments, analogs or homologs thereof, that contain the antibody derived binding domain, are utilized as pharmacologically-active compounds [hereinafter "Therapeutics"].

An anti-ORFX antibody (e.g., monoclonal antibody) can be used to isolate ORFX by standard techniques, such as affinity chromatography or immunoprecipitation. An anti-ORFX antibody can facilitate the purification of natural ORFX from cells and of recombinantly produced ORFX expressed in host cells. Moreover, an anti-ORFX antibody can be used to detect ORFX protein (e.g., in a cellular lysate or cell supernatant) in order to evaluate the abundance and pattern of expression of the ORFX protein. Anti-ORFX antibodies can be used diagnostically to monitor protein levels in tissue as part of a clinical testing procedure, e.g., to, for example, determine the efficacy of a given treatment regimen. Detection can be facilitated by coupling (i.e., physically linking) the antibody to a detectable substance. Examples of detectable substances include various enzymes, prosthetic groups, fluorescent materials, luminescent materials, bioluminescent materials, and radioactive materials. Examples of suitable enzymes include horseradish peroxidase, alkaline phosphatase, β-galactosidase, or acetylcholinesterase; examples of suitable prosthetic group complexes include streptavidin/biotin and avidin/biotin; examples of suitable fluorescent materials include umbelliferone, fluorescein, fluorescein isothiocyanate, rhodamine, dichlorotriazinylamine fluorescein, dansyl chloride or

phycoerythrin; an example of a luminescent material includes luminol; examples of bioluminescent materials include luciferase, luciferin, and aequorin, and examples of suitable radioactive material include <sup>125</sup>I, <sup>131</sup>I, <sup>35</sup>S or <sup>3</sup>H.

## **ORFX** Recombinant Vectors and Host Cells

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Another aspect of the invention pertains to vectors, preferably expression vectors, containing a nucleic acid encoding ORFX protein, or derivatives, fragments, analogs or homologs thereof. As used herein, the term "vector" refers to a nucleic acid molecule capable of transporting another nucleic acid to which it has been linked. One type of vector is a "plasmid", which refers to a circular double stranded DNA loop into which additional DNA segments can be ligated. Another type of vector is a viral vector, wherein additional DNA segments can be ligated into the viral genome. Certain vectors are capable of autonomous replication in a host cell into which they are introduced (e.g., bacterial vectors having a bacterial origin of replication and episomal mammalian vectors). Other vectors (e.g., non-episomal mammalian vectors) are integrated into the genome of a host cell upon introduction into the host cell, and thereby are replicated along with the host genome. Moreover, certain vectors are capable of directing the expression of genes to which they are operatively linked. Such vectors are referred to herein as "expression vectors". In general, expression vectors of utility in recombinant DNA techniques are often in the form of plasmids. In the present specification, "plasmid" and "vector" can be used interchangeably as the plasmid is the most commonly used form of vector. However, the invention is intended to include such other forms of expression vectors, such as viral vectors (e.g., replication defective retroviruses, adenoviruses and adeno-associated viruses), which serve equivalent functions.

The recombinant expression vectors of the invention comprise a nucleic acid of the invention in a form suitable for expression of the nucleic acid in a host cell, which means that the recombinant expression vectors include one or more regulatory sequences, selected on the basis of the host cells to be used for expression, that is operatively linked to the nucleic acid sequence to be expressed. Within a recombinant expression vector, "operably linked" is intended to mean that the nucleotide sequence of interest is linked to the regulatory sequence(s) in a manner that allows for expression of the nucleotide sequence (e.g., in an in vitro transcription/translation system or in a host cell when the vector is introduced into the host cell). The term "regulatory sequence" is intended to includes promoters, enhancers and other expression control elements

(e.g., polyadenylation signals). Such regulatory sequences are described, for example, in Goeddel; GENE EXPRESSION TECHNOLOGY: METHODS IN ENZYMOLOGY 185, Academic Press, San Diego, Calif. (1990). Regulatory sequences include those that direct constitutive expression of a nucleotide sequence in many types of host cell and those that direct expression of the nucleotide sequence only in certain host cells (e.g., tissue-specific regulatory sequences). It will be appreciated by those skilled in the art that the design of the expression vector can depend on such factors as the choice of the host cell to be transformed, the level of expression of protein desired, etc. The expression vectors of the invention can be introduced into host cells to thereby produce proteins or peptides, including fusion proteins or peptides, encoded by nucleic acids as described herein (e.g., ORFX proteins, mutant forms of ORFX, fusion proteins, etc.).

The recombinant expression vectors of the invention can be designed for expression of ORFX in prokaryotic or eukaryotic cells. For example, ORFX can be expressed in bacterial cells such as *E. coli*, insect cells (using baculovirus expression vectors) yeast cells or mammalian cells. Suitable host cells are discussed further in Goeddel, GENE EXPRESSION TECHNOLOGY: METHODS IN ENZYMOLOGY 185, Academic Press, San Diego, Calif. (1990). Alternatively, the recombinant expression vector can be transcribed and translated *in vitro*, for example using T7 promoter regulatory sequences and T7 polymerase.

Expression of proteins in prokaryotes is most often carried out in *E. coli* with vectors containing constitutive or inducible promoters directing the expression of either fusion or non-fusion proteins. Fusion vectors add a number of amino acids to a protein encoded therein, usually to the amino terminus of the recombinant protein. Such fusion vectors typically serve three purposes: (1) to increase expression of recombinant protein; (2) to increase the solubility of the recombinant protein; and (3) to aid in the purification of the recombinant protein by acting as a ligand in affinity purification. Often, in fusion expression vectors, a proteolytic cleavage site is introduced at the junction of the fusion moiety and the recombinant protein to enable separation of the recombinant protein from the fusion moiety subsequent to purification of the fusion protein. Such enzymes, and their cognate recognition sequences, include Factor Xa, thrombin and enterokinase. Typical fusion expression vectors include pGEX (Pharmacia Biotech Inc; Smith and Johnson (1988) *Gene* 67:31-40), pMAL (New England Biolabs, Beverly, Mass.) and pRIT5 (Pharmacia, Piscataway, N.J.) that fuse glutathione S-transferase (GST), maltose E binding protein, or protein A, respectively, to the target recombinant protein.

Examples of suitable inducible non-fusion *E. coli* expression vectors include pTrc (Amrann *et al.*, (1988) *Gene* 69:301-315) and pET 11d (Studier *et al.*, GENE EXPRESSION TECHNOLOGY: METHODS IN ENZYMOLOGY 185, Academic Press, San Diego, Calif. (1990) 60-89).

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One strategy to maximize recombinant protein expression in *E. coli* is to express the protein in a host bacteria with an impaired capacity to proteolytically cleave the recombinant protein. See, Gottesman, GENE EXPRESSION TECHNOLOGY: METHODS IN ENZYMOLOGY 185, Academic Press, San Diego, Calif. (1990) 119-128. Another strategy is to alter the nucleic acid sequence of the nucleic acid to be inserted into an expression vector so that the individual codons for each amino acid are those preferentially utilized in *E. coli* (Wada *et al.*, (1992) *Nucleic Acids Res.* 20:2111-2118). Such alteration of nucleic acid sequences of the invention can be carried out by standard DNA synthesis techniques.

In another embodiment, the ORFX expression vector is a yeast expression vector. Examples of vectors for expression in yeast *S. cerivisae* include pYepSec1 (Baldari, *et al.*, (1987) *EMBO J* 6:229-234), pMFa (Kurjan and Herskowitz, (1982) *Cell* 30:933-943), pJRY88 (Schultz *et al.*, (1987) *Gene* 54:113-123), pYES2 (Invitrogen Corporation, San Diego, Calif.), and picZ (InVitrogen Corp, San Diego, Calif.).

Alternatively, ORFX can be expressed in insect cells using baculovirus expression vectors. Baculovirus vectors available for expression of proteins in cultured insect cells (e.g., SF9 cells) include the pAc series (Smith et al. (1983) Mol Cell Biol 3:2156-2165) and the pVL series (Lucklow and Summers (1989) Virology 170:31-39).

In yet another embodiment, a nucleic acid of the invention is expressed in mammalian cells using a mammalian expression vector. Examples of mammalian expression vectors include pCDM8 (Seed (1987) Nature 329:840) and pMT2PC (Kaufman et al. (1987) EMBO J 6: 187-195). When used in mammalian cells, the expression vector's control functions are often provided by viral regulatory elements. For example, commonly used promoters are derived from polyoma, Adenovirus 2, cytomegalovirus and Simian Virus 40. For other suitable expression systems for both prokaryotic and eukaryotic cells. See, e.g., Chapters 16 and 17 of Sambrook et al., MOLECULAR CLONING: A LABORATORY MANUAL. 2nd ed., Cold Spring Harbor Laboratory, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., 1989.

In another embodiment, the recombinant mammalian expression vector is capable of directing expression of the nucleic acid preferentially in a particular cell type (e.g., tissue-specific regulatory elements are used to express the nucleic acid). Tissue-specific regulatory elements are known in the art. Non-limiting examples of suitable tissue-specific promoters include the albumin promoter (liver-specific; Pinkert et al. (1987) Genes Dev 1:268-277), lymphoid-specific promoters (Calame and Eaton (1988) Adv Immunol 43:235-275), in particular promoters of T cell receptors (Winoto and Baltimore (1989) EMBO J 8:729-733) and immunoglobulins (Banerji et al. (1983) Cell 33:729-740; Queen and Baltimore (1983) Cell 33:741-748), neuron-specific promoters (e.g., the neurofilament promoter; Byrne and Ruddle (1989) PNAS 86:5473-5477), pancreas-specific promoters (Edlund et al. (1985) Science 230:912-916), and mammary gland-specific promoters (e.g., milk whey promoter; U.S. Pat. No. 4,873,316 and European Application Publication No. 264,166). Developmentally-regulated promoters are also encompassed, e.g., the murine hox promoters (Kessel and Gruss (1990) Science 249:374-379) and the α-fetoprotein promoter (Campes and Tilghman (1989) Genes Dev 3:537-546).

The invention further provides a recombinant expression vector comprising a DNA molecule of the invention cloned into the expression vector in an antisense orientation. That is, the DNA molecule is operatively linked to a regulatory sequence in a manner that allows for expression (by transcription of the DNA molecule) of an RNA molecule that is antisense to ORFX mRNA. Regulatory sequences operatively linked to a nucleic acid cloned in the antisense orientation can be chosen that direct the continuous expression of the antisense RNA molecule in a variety of cell types, for instance viral promoters and/or enhancers, or regulatory sequences can be chosen that direct constitutive, tissue specific or cell type specific expression of antisense RNA. The antisense expression vector can be in the form of a recombinant plasmid, phagemid or attenuated virus in which antisense nucleic acids are produced under the control of a high efficiency regulatory region, the activity of which can be determined by the cell type into which the vector is introduced. For a discussion of the regulation of gene expression using antisense genes see Weintraub *et al.*, "Antisense RNA as a molecular tool for genetic analysis," Reviews—Trends in Genetics, Vol. 1(1) 1986.

Another aspect of the invention pertains to host cells into which a recombinant expression vector of the invention has been introduced. The terms "host cell" and "recombinant

host cell" are used interchangeably herein. It is understood that such terms refer not only to the particular subject cell but to the progeny or potential progeny of such a cell. Because certain modifications may occur in succeeding generations due to either mutation or environmental influences, such progeny may not, in fact, be identical to the parent cell, but are still included within the scope of the term as used herein.

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A host cell can be any prokaryotic or eukaryotic cell. For example, ORFX protein can be expressed in bacterial cells such as *E. coli*, insect cells, yeast or mammalian cells (such as Chinese hamster ovary cells (CHO) or COS cells). Other suitable host cells are known to those skilled in the art.

Vector DNA can be introduced into prokaryotic or eukaryotic cells via conventional transformation or transfection techniques. As used herein, the terms "transformation" and "transfection" are intended to refer to a variety of art-recognized techniques for introducing foreign nucleic acid (e.g., DNA) into a host cell, including calcium phosphate or calcium chloride co-precipitation, DEAE-dextran-mediated transfection, lipofection, or electroporation. Suitable methods for transforming or transfecting host cells can be found in Sambrook, et al. (MOLECULAR CLONING: A LABORATORY MANUAL. 2nd ed., Cold Spring Harbor Laboratory, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., 1989), and other laboratory manuals.

For stable transfection of mammalian cells, it is known that, depending upon the expression vector and transfection technique used, only a small fraction of cells may integrate the foreign DNA into their genome. In order to identify and select these integrants, a gene that encodes a selectable marker (e.g., resistance to antibiotics) is generally introduced into the host cells along with the gene of interest. Various selectable markers include those that confer resistance to drugs, such as G418, hygromycin and methotrexate. Nucleic acid encoding a selectable marker can be introduced into a host cell on the same vector as that encoding ORFX or can be introduced on a separate vector. Cells stably transfected with the introduced nucleic acid can be identified by drug selection (e.g., cells that have incorporated the selectable marker gene will survive, while the other cells die).

A host cell of the invention, such as a prokaryotic or eukaryotic host cell in culture, can be used to produce (i.e., express) ORFX protein. Accordingly, the invention further provides methods for producing ORFX protein using the host cells of the invention. In one embodiment,

the method comprises culturing the host cell of invention (into which a recombinant expression vector encoding ORFX has been introduced) in a suitable medium such that ORFX protein is produced. In another embodiment, the method further comprises isolating ORFX from the medium or the host cell.

## Transgenic animals

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The host cells of the invention can also be used to produce nonhuman transgenic animals. For example, in one embodiment, a host cell of the invention is a fertilized oocyte or an embryonic stem cell into which ORFX-coding sequences have been introduced. Such host cells can then be used to create non-human transgenic animals in which exogenous ORFX sequences have been introduced into their genome or homologous recombinant animals in which endogenous ORFX sequences have been altered. Such animals are useful for studying the function and/or activity of ORFX and for identifying and/or evaluating modulators of ORFX activity. As used herein, a "transgenic animal" is a non-human animal, preferably a mammal, more preferably a rodent such as a rat or mouse, in which one or more of the cells of the animal includes a transgene. Other examples of transgenic animals include non-human primates, sheep, dogs, cows, goats, chickens, amphibians, etc. A transgene is exogenous DNA that is integrated into the genome of a cell from which a transgenic animal develops and that remains in the genome of the mature animal, thereby directing the expression of an encoded gene product in one or more cell types or tissues of the transgenic animal. As used herein, a "homologous recombinant animal" is a non-human animal, preferably a mammal, more preferably a mouse, in which an endogenous ORFX gene has been altered by homologous recombination between the endogenous gene and an exogenous DNA molecule introduced into a cell of the animal, e.g., an embryonic cell of the animal, prior to development of the animal.

A transgenic animal of the invention can be created by introducing ORFX-encoding nucleic acid into the male pronuclei of a fertilized oocyte, e.g., by microinjection, retroviral infection, and allowing the oocyte to develop in a pseudopregnant female foster animal. The human ORFX DNA sequence of SEQ ID NO:2n-1 (wherein n = 1 to 3161) can be introduced as a transgene into the genome of a non-human animal. Alternatively, a nonhuman homologue of the human ORFX gene, such as a mouse ORFX gene, can be isolated based on hybridization to the human ORFX cDNA (described further above) and used as a transgene. Intronic sequences and polyadenylation signals can also be included in the transgene to increase the efficiency of

expression of the transgene. A tissue-specific regulatory sequence(s) can be operably linked to the ORFX transgene to direct expression of ORFX protein to particular cells. Methods for generating transgenic animals via embryo manipulation and microinjection, particularly animals such as mice, have become conventional in the art and are described, for example, in U.S. Pat. Nos. 4,736,866; 4,870,009; and 4,873,191; and Hogan 1986, In: MANIPULATING THE MOUSE EMBRYO, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y. Similar methods are used for production of other transgenic animals. A transgenic founder animal can be identified based upon the presence of the ORFX transgene in its genome and/or expression of ORFX mRNA in tissues or cells of the animals. A transgenic founder animal can then be used to breed additional animals carrying the transgene. Moreover, transgenic animals carrying a transgene encoding ORFX can further be bred to other transgenic animals carrying other transgenes.

To create a homologous recombinant animal, a vector is prepared which contains at least a portion of a ORFX gene into which a deletion, addition or substitution has been introduced to thereby alter, e.g., functionally disrupt, the ORFX gene. The ORFX gene can be a human gene (e.g., SEQ ID NO:2n-1 (wherein n = 1 to 3161)), but more preferably, is a non-human homologue of a human ORFX gene. For example, a mouse homologue of human ORFX gene of SEQ ID NO:2n-1 (wherein n = 1 to 3161) can be used to construct a homologous recombination vector suitable for altering an endogenous ORFX gene in the mouse genome. In one embodiment, the vector is designed such that, upon homologous recombination, the endogenous ORFX gene is functionally disrupted (i.e., no longer encodes a functional protein; also referred to as a "knock out" vector).

Alternatively, the vector can be designed such that, upon homologous recombination, the endogenous ORFX gene is mutated or otherwise altered but still encodes functional protein (e.g., the upstream regulatory region can be altered to thereby alter the expression of the endogenous ORFX protein). In the homologous recombination vector, the altered portion of the ORFX gene is flanked at its 5' and 3' ends by additional nucleic acid of the ORFX gene to allow for homologous recombination to occur between the exogenous ORFX gene carried by the vector and an endogenous ORFX gene in an embryonic stem cell. The additional flanking ORFX nucleic acid is of sufficient length for successful homologous recombination with the endogenous gene. Typically, several kilobases of flanking DNA (both at the 5' and 3' ends) are included in the vector. See e.g., Thomas et al. (1987) Cell 51:503 for a description of

homologous recombination vectors. The vector is introduced into an embryonic stem cell line (e.g., by electroporation) and cells in which the introduced ORFX gene has homologously recombined with the endogenous ORFX gene are selected (see e.g., Li et al. (1992) Cell 69:915).

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The selected cells are then injected into a blastocyst of an animal (e.g., a mouse) to form aggregation chimeras. See e.g., Bradley 1987, In: TERATOCARCINOMAS AND EMBRYONIC STEM CELLS: A PRACTICAL APPROACH, Robertson, ed. IRL, Oxford, pp. 113-152. A chimeric embryo can then be implanted into a suitable pseudopregnant female foster animal and the embryo brought to term. Progeny harboring the homologously recombined DNA in their germ cells can be used to breed animals in which all cells of the animal contain the homologously recombined DNA by germline transmission of the transgene. Methods for constructing homologous recombination vectors and homologous recombinant animals are described further in Bradley (1991) Curr Opin Biotechnol 2:823-829; PCT International Publication Nos.: WO 90/11354; WO 91/01140; WO 92/0968; and WO 93/04169.

In another embodiment, transgenic non-humans animals can be produced that contain selected systems that allow for regulated expression of the transgene. One example of such a system is the cre/loxP recombinase system of bacteriophage P1. For a description of the cre/loxP recombinase system, see, e.g., Lakso et al. (1992) PNAS 89:6232-6236. Another example of a recombinase system is the FLP recombinase system of Saccharomyces cerevisiae (O'Gorman et al. (1991) Science 251:1351-1355. If a cre/loxP recombinase system is used to regulate expression of the transgene, animals containing transgenes encoding both the Cre recombinase and a selected protein are required. Such animals can be provided through the construction of "double" transgenic animals, e.g., by mating two transgenic animals, one containing a transgene encoding a selected protein and the other containing a transgene encoding a recombinase.

Clones of the non-human transgenic animals described herein can also be produced according to the methods described in Wilmut et al. (1997) Nature 385:810-813. In brief, a cell, e.g., a somatic cell, from the transgenic animal can be isolated and induced to exit the growth cycle and enter G<sub>0</sub> phase. The quiescent cell can then be fused, e.g., through the use of electrical pulses, to an enucleated oocyte from an animal of the same species from which the quiescent cell is isolated. The reconstructed oocyte is then cultured such that it develops to morula or blastocyte and then transferred to pseudopregnant female foster animal. The offspring borne of

this female foster animal will be a clone of the animal from which the cell, e.g., the somatic cell, is isolated.

## Pharmaceutical Compositions

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The ORFX nucleic acid molecules, ORFX proteins, and anti-ORFX antibodies (also referred to herein as "active compounds") of the invention, and derivatives, fragments, analogs and homologs thereof, can be incorporated into pharmaceutical compositions suitable for administration. Such compositions typically comprise the nucleic acid molecule, protein, or antibody and a pharmaceutically acceptable carrier. As used herein, "pharmaceutically acceptable carrier" is intended to include any and all solvents, dispersion media, coatings, antibacterial and antifungal agents, isotonic and absorption delaying agents, and the like, compatible with pharmaceutical administration. Suitable carriers are described in the most recent edition of Remington's Pharmaceutical Sciences, a standard reference text in the field, which is incorporated herein by reference. Preferred examples of such carriers or diluents include, but are not limited to, water, saline, finger's solutions, dextrose solution, and 5% human serum albumin. Liposomes and non-aqueous vehicles such as fixed oils may also be used. The use of such media and agents for pharmaceutically active substances is well known in the art. Except insofar as any conventional media or agent is incompatible with the active compound, use thereof in the compositions is contemplated. Supplementary active compounds can also be incorporated into the compositions.

A pharmaceutical composition of the invention is formulated to be compatible with its intended route of administration. Examples of routes of administration include parenteral, e.g., intravenous, intradermal, subcutaneous, oral (e.g., inhalation), transdermal (topical), transmucosal, and rectal administration. Solutions or suspensions used for parenteral, intradermal, or subcutaneous application can include the following components: a sterile diluent such as water for injection, saline solution, fixed oils, polyethylene glycols, glycerine, propylene glycol or other synthetic solvents; antibacterial agents such as benzyl alcohol or methyl parabens; antioxidants such as ascorbic acid or sodium bisulfite; chelating agents such as ethylenediaminetetraacetic acid; buffers such as acetates, citrates or phosphates, and agents for the adjustment of tonicity such as sodium chloride or dextrose. The pH can be adjusted with acids or bases, such as hydrochloric acid or sodium hydroxide. The parenteral preparation can be enclosed in ampoules, disposable syringes or multiple dose vials made of glass or plastic.

Pharmaceutical compositions suitable for injectable use include sterile aqueous solutions (where water soluble) or dispersions and sterile powders for the extemporaneous preparation of sterile injectable solutions or dispersion. For intravenous administration, suitable carriers include physiological saline, bacteriostatic water, Cremophor EL<sup>TM</sup> (BASF, Parsippany, N.J.) or phosphate buffered saline (PBS). In all cases, the composition must be sterile and should be fluid to the extent that easy syringeability exists. It must be stable under the conditions of manufacture and storage and must be preserved against the contaminating action of microorganisms such as bacteria and fungi. The carrier can be a solvent or dispersion medium containing, for example, water, ethanol, polyol (for example, glycerol, propylene glycol, and liquid polyethylene glycol, and the like), and suitable mixtures thereof. The proper fluidity can be maintained, for example, by the use of a coating such as lecithin, by the maintenance of the required particle size in the case of dispersion and by the use of surfactants. Prevention of the action of microorganisms can be achieved by various antibacterial and antifungal agents, for example, parabens, chlorobutanol, phenol, ascorbic acid, thimerosal, and the like. In many cases, it will be preferable to include isotonic agents, for example, sugars, polyalcohols such as manitol, sorbitol, sodium chloride in the composition. Prolonged absorption of the injectable compositions can be brought about by including in the composition an agent which delays absorption, for example, aluminum monostearate and gelatin.

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Sterile injectable solutions can be prepared by incorporating the active compound (e.g., a ORFX protein or anti-ORFX antibody) in the required amount in an appropriate solvent with one or a combination of ingredients enumerated above, as required, followed by filtered sterilization. Generally, dispersions are prepared by incorporating the active compound into a sterile vehicle that contains a basic dispersion medium and the required other ingredients from those enumerated above. In the case of sterile powders for the preparation of sterile injectable solutions, methods of preparation are vacuum drying and freeze-drying that yields a powder of the active ingredient plus any additional desired ingredient from a previously sterile-filtered solution thereof.

Oral compositions generally include an inert diluent or an edible carrier. They can be enclosed in gelatin capsules or compressed into tablets. For the purpose of oral therapeutic administration, the active compound can be incorporated with excipients and used in the form of tablets, troches, or capsules. Oral compositions can also be prepared using a fluid carrier for use

as a mouthwash, wherein the compound in the fluid carrier is applied orally and swished and expectorated or swallowed. Pharmaceutically compatible binding agents, and/or adjuvant materials can be included as part of the composition. The tablets, pills, capsules, troches and the like can contain any of the following ingredients, or compounds of a similar nature: a binder such as microcrystalline cellulose, gum tragacanth or gelatin; an excipient such as starch or lactose, a disintegrating agent such as alginic acid, Primogel, or corn starch; a lubricant such as magnesium stearate or Sterotes; a glidant such as colloidal silicon dioxide; a sweetening agent such as sucrose or saccharin; or a flavoring agent such as peppermint, methyl salicylate, or orange flavoring.

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For administration by inhalation, the compounds are delivered in the form of an aerosol spray from pressured container or dispenser which contains a suitable propellant, e.g., a gas such as carbon dioxide, or a nebulizer.

Systemic administration can also be by transmucosal or transdermal means. For transmucosal or transdermal administration, penetrants appropriate to the barrier to be permeated are used in the formulation. Such penetrants are generally known in the art, and include, for example, for transmucosal administration, detergents, bile salts, and fusidic acid derivatives. Transmucosal administration can be accomplished through the use of nasal sprays or suppositories. For transdermal administration, the active compounds are formulated into ointments, salves, gels, or creams as generally known in the art.

The compounds can also be prepared in the form of suppositories (e.g., with conventional suppository bases such as cocoa butter and other glycerides) or retention enemas for rectal delivery.

In one embodiment, the active compounds are prepared with carriers that will protect the compound against rapid elimination from the body, such as a controlled release formulation, including implants and microencapsulated delivery systems. Biodegradable, biocompatible polymers can be used, such as ethylene vinyl acetate, polyanhydrides, polyglycolic acid, collagen, polyorthoesters, and polylactic acid. Methods for preparation of such formulations will be apparent to those skilled in the art. The materials can also be obtained commercially from Alza Corporation and Nova Pharmaceuticals, Inc. Liposomal suspensions (including liposomes targeted to infected cells with monoclonal antibodies to viral antigens) can also be used as pharmaceutically acceptable carriers. These can be prepared according to methods known to those skilled in the art, for example, as described in U.S. Pat. No. 4,522,811.

It is especially advantageous to formulate oral or parenteral compositions in dosage unit form for ease of administration and uniformity of dosage. Dosage unit form as used herein refers to physically discrete units suited as unitary dosages for the subject to be treated; each unit containing a predetermined quantity of active compound calculated to produce the desired therapeutic effect in association with the required pharmaceutical carrier. The specification for the dosage unit forms of the invention are dictated by and directly dependent on the unique characteristics of the active compound and the particular therapeutic effect to be achieved.

The nucleic acid molecules of the invention can be inserted into vectors and used as gene therapy vectors. Gene therapy vectors can be delivered to a subject by any of a number of routes, e.g., as described in U.S. Patent Nos. 5,703,055. Delivery can thus also include, e.g., intravenous injection, local administration (see U.S. Pat. No. 5,328,470) or stereotactic injection (see e.g., Chen et al. (1994) PNAS 91:3054-3057). The pharmaceutical preparation of the gene therapy vector can include the gene therapy vector in an acceptable diluent, or can comprise a slow release matrix in which the gene delivery vehicle is imbedded. Alternatively, where the complete gene delivery vector can be produced intact from recombinant cells, e.g., retroviral vectors, the pharmaceutical preparation can include one or more cells that produce the gene delivery system.

The pharmaceutical compositions can be included in a container, pack, or dispenser together with instructions for administration.

### Additional Uses and Methods of the Invention

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The nucleic acid molecules, proteins, protein homologues, and antibodies described herein can be used in one or more of the following methods: (a) screening assays; (b) detection assays (e.g., chromosomal mapping, cell and tissue typing, forensic biology), (c) predictive medicine (e.g., diagnostic assays, prognostic assays, monitoring clinical trials, and pharmacogenomics); and (d) methods of treatment (e.g., therapeutic and prophylactic).

The isolated nucleic acid molecules of the invention can be used to express ORFX protein (e.g., via a recombinant expression vector in a host cell in gene therapy applications), to detect ORFX mRNA (e.g., in a biological sample) or a genetic lesion in a ORFX gene, and to modulate ORFX activity, as described further below. In addition, the ORFX proteins can be used to screen drugs or compounds that modulate the ORFX activity or expression as well as to treat disorders characterized by insufficient or excessive production of ORFX protein, for

example proliferative or differentiative disorders, or production of ORFX protein forms that have decreased or aberrant activity compared to ORFX wild type protein. In addition, the anti-ORFX antibodies of the invention can be used to detect and isolate ORFX proteins and modulate ORFX activity.

This invention further pertains to novel agents identified by the above described screening assays and uses thereof for treatments as described herein.

### Screening Assays

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The invention provides a method (also referred to herein as a "screening assay") for identifying modulators, *i.e.*, candidate or test compounds or agents (*e.g.*, peptides, peptidomimetics, small molecules or other drugs) that bind to ORFX proteins or have a stimulatory or inhibitory effect on, for example, ORFX expression or ORFX activity.

In one embodiment, the invention provides assays for screening candidate or test compounds which bind to or modulate the activity of a ORFX protein or polypeptide or biologically active portion thereof. The test compounds of the present invention can be obtained using any of the numerous approaches in combinatorial library methods known in the art, including: biological libraries; spatially addressable parallel solid phase or solution phase libraries; synthetic library methods requiring deconvolution; the "one-bead one-compound" library method; and synthetic library methods using affinity chromatography selection. The biological library approach is limited to peptide libraries, while the other four approaches are applicable to peptide, non-peptide oligomer or small molecule libraries of compounds (Lam (1997) Anticancer Drug Des 12:145).

Examples of methods for the synthesis of molecular libraries can be found in the art, for example in: DeWitt et al. (1993) Proc Natl Acad Sci U.S.A. 90:6909; Erb et al. (1994) Proc Natl Acad Sci U.S.A. 91:11422; Zuckermann et al. (1994) J Med Chem 37:2678; Cho et al. (1993) Science 261:1303; Carrell et al. (1994) Angew Chem Int Ed Engl 33:2059; Carell et al. (1994) Angew Chem Int Ed Engl 33:2061; and Gallop et al. (1994) J Med Chem 37:1233.

Libraries of compounds may be presented in solution (e.g., Houghten (1992)

Biotechniques 13:412-421), or on beads (Lam (1991) Nature 354:82-84), on chips (Fodor (1993)

Nature 364:555-556), bacteria (Ladner U.S. Pat. No. 5,223,409), spores (Ladner USP '409),

plasmids (Cull et al. (1992) Proc Natl Acad Sci USA 89:1865-1869) or on phage (Scott and

Smith (1990) Science 249:386-390; Devlin (1990) Science 249:404-406; Cwirla et al. (1990) Proc Natl Acad Sci U.S.A. 87:6378-6382; Felici (1991) J Mol Biol 222:301-310; Ladner above.).

In one embodiment, an assay is a cell-based assay in which a cell which expresses a membrane-bound form of ORFX protein, or a biologically active portion thereof, on the cell surface is contacted with a test compound and the ability of the test compound to bind to a ORFX protein determined. The cell, for example, can of mammalian origin or a yeast cell. Determining the ability of the test compound to bind to the ORFX protein can be accomplished, for example, by coupling the test compound with a radioisotope or enzymatic label such that binding of the test compound to the ORFX protein or biologically active portion thereof can be determined by detecting the labeled compound in a complex. For example, test compounds can be labeled with <sup>125</sup>I, <sup>35</sup>S, <sup>14</sup>C, or <sup>3</sup>H, either directly or indirectly, and the radioisotope detected by direct counting of radioemission or by scintillation counting. Alternatively, test compounds can be enzymatically labeled with, for example, horseradish peroxidase, alkaline phosphatase, or luciferase, and the enzymatic label detected by determination of conversion of an appropriate substrate to product. In one embodiment, the assay comprises contacting a cell which expresses a membrane-bound form of ORFX protein, or a biologically active portion thereof, on the cell surface with a known compound which binds ORFX to form an assay mixture, contacting the assay mixture with a test compound, and determining the ability of the test compound to interact with a ORFX protein, wherein determining the ability of the test compound to interact with a ORFX protein comprises determining the ability of the test compound to preferentially bind to ORFX or a biologically active portion thereof as compared to the known compound.

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In another embodiment, an assay is a cell-based assay comprising contacting a cell expressing a membrane-bound form of ORFX protein, or a biologically active portion thereof, on the cell surface with a test compound and determining the ability of the test compound to modulate (e.g., stimulate or inhibit) the activity of the ORFX protein or biologically active portion thereof. Determining the ability of the test compound to modulate the activity of ORFX or a biologically active portion thereof can be accomplished, for example, by determining the ability of the ORFX protein to bind to or interact with a ORFX target molecule. As used herein, a "target molecule" is a molecule with which a ORFX protein binds or interacts in nature, for example, a molecule on the surface of a cell which expresses a ORFX interacting protein, a molecule on the surface of a second cell, a molecule in the extracellular milieu, a molecule

associated with the internal surface of a cell membrane or a cytoplasmic molecule. A ORFX target molecule can be a non-ORFX molecule or a ORFX protein or polypeptide of the present invention. In one embodiment, a ORFX target molecule is a component of a signal transduction pathway that facilitates transduction of an extracellular signal (e.g., a signal generated by binding of a compound to a membrane-bound ORFX molecule) through the cell membrane and into the cell. The target, for example, can be a second intercellular protein that has catalytic activity or a protein that facilitates the association of downstream signaling molecules with ORFX.

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Determining the ability of the ORFX protein to bind to or interact with a ORFX target molecule can be accomplished by one of the methods described above for determining direct binding. In one embodiment, determining the ability of the ORFX protein to bind to or interact with a ORFX target molecule can be accomplished by determining the activity of the target molecule. For example, the activity of the target molecule can be determined by detecting induction of a cellular second messenger of the target (*i.e.* intracellular Ca<sup>2+</sup>, diacylglycerol, IP<sub>3</sub>, etc.), detecting catalytic/enzymatic activity of the target an appropriate substrate, detecting the induction of a reporter gene (comprising a ORFX-responsive regulatory element operatively linked to a nucleic acid encoding a detectable marker, *e.g.*, luciferase), or detecting a cellular response, for example, cell survival, cellular differentiation, or cell proliferation.

In yet another embodiment, an assay of the present invention is a cell-free assay comprising contacting a ORFX protein or biologically active portion thereof with a test compound and determining the ability of the test compound to bind to the ORFX protein or biologically active portion thereof. Binding of the test compound to the ORFX protein can be determined either directly or indirectly as described above. In one embodiment, the assay comprises contacting the ORFX protein or biologically active portion thereof with a known compound which binds ORFX to form an assay mixture, contacting the assay mixture with a test compound, and determining the ability of the test compound to interact with a ORFX protein, wherein determining the ability of the test compound to interact with a ORFX protein comprises determining the ability of the test compound to preferentially bind to ORFX or biologically active portion thereof as compared to the known compound.

In another embodiment, an assay is a cell-free assay comprising contacting ORFX protein or biologically active portion thereof with a test compound and determining the ability of the test compound to modulate (e.g., stimulate or inhibit) the activity of the ORFX protein or

biologically active portion thereof. Determining the ability of the test compound to modulate the activity of ORFX can be accomplished, for example, by determining the ability of the ORFX protein to bind to a ORFX target molecule by one of the methods described above for determining direct binding. In an alternative embodiment, determining the ability of the test compound to modulate the activity of ORFX can be accomplished by determining the ability of the ORFX protein further modulate a ORFX target molecule. For example, the catalytic/enzymatic activity of the target molecule on an appropriate substrate can be determined as previously described.

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In yet another embodiment, the cell-free assay comprises contacting the ORFX protein or biologically active portion thereof with a known compound which binds ORFX to form an assay mixture, contacting the assay mixture with a test compound, and determining the ability of the test compound to interact with a ORFX protein, wherein determining the ability of the test compound to interact with a ORFX protein comprises determining the ability of the ORFX protein to preferentially bind to or modulate the activity of a ORFX target molecule.

The cell-free assays of the present invention are amenable to use of both the soluble form or the membrane-bound form of ORFX. In the case of cell-free assays comprising the membrane-bound form of ORFX, it may be desirable to utilize a solubilizing agent such that the membrane-bound form of ORFX is maintained in solution. Examples of such solubilizing agents include non-ionic detergents such as n-octylglucoside, n-dodecylglucoside, n-dodecylmaltoside, octanoyl-N-methylglucamide, decanoyl-N-methylglucamide, Triton<sup>®</sup> X-100, Triton<sup>®</sup> X-114, Thesit<sup>®</sup>, Isotridecypoly(ethylene glycol ether)<sub>n</sub>, N-dodecyl--N,N-dimethyl-3-ammonio-1-propane sulfonate, 3-(3-cholamidopropyl)dimethylamminiol-1-propane sulfonate (CHAPSO).

In more than one embodiment of the above assay methods of the present invention, it may be desirable to immobilize either ORFX or its target molecule to facilitate separation of complexed from uncomplexed forms of one or both of the proteins, as well as to accommodate automation of the assay. Binding of a test compound to ORFX, or interaction of ORFX with a target molecule in the presence and absence of a candidate compound, can be accomplished in any vessel suitable for containing the reactants. Examples of such vessels include microtiter plates, test tubes, and micro-centrifuge tubes. In one embodiment, a fusion protein can be provided that adds a domain that allows one or both of the proteins to be bound to a matrix. For

example, GST-ORFX fusion proteins or GST-target fusion proteins can be adsorbed onto glutathione sepharose beads (Sigma Chemical, St. Louis, MO) or glutathione derivatized microtiter plates, that are then combined with the test compound or the test compound and either the non-adsorbed target protein or ORFX protein, and the mixture is incubated under conditions conducive to complex formation (e.g., at physiological conditions for salt and pH). Following incubation, the beads or microtiter plate wells are washed to remove any unbound components, the matrix immobilized in the case of beads, complex determined either directly or indirectly, for example, as described above. Alternatively, the complexes can be dissociated from the matrix, and the level of ORFX binding or activity determined using standard techniques.

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Other techniques for immobilizing proteins on matrices can also be used in the screening assays of the invention. For example, either ORFX or its target molecule can be immobilized utilizing conjugation of biotin and streptavidin. Biotinylated ORFX or target molecules can be prepared from biotin-NHS (N-hydroxy-succinimide) using techniques well known in the art (e.g., biotinylation kit, Pierce Chemicals, Rockford, Ill.), and immobilized in the wells of streptavidin-coated 96 well plates (Pierce Chemical). Alternatively, antibodies reactive with ORFX or target molecules, but which do not interfere with binding of the ORFX protein to its target molecule, can be derivatized to the wells of the plate, and unbound target or ORFX trapped in the wells by antibody conjugation. Methods for detecting such complexes, in addition to those described above for the GST-immobilized complexes, include immunodetection of complexes using antibodies reactive with the ORFX or target molecule, as well as enzyme-linked assays that rely on detecting an enzymatic activity associated with the ORFX or target molecule.

In another embodiment, modulators of ORFX expression are identified in a method wherein a cell is contacted with a candidate compound and the expression of ORFX mRNA or protein in the cell is determined. The level of expression of ORFX mRNA or protein in the presence of the candidate compound is compared to the level of expression of ORFX mRNA or protein in the absence of the candidate compound. The candidate compound can then be identified as a modulator of ORFX expression based on this comparison. For example, when expression of ORFX mRNA or protein is greater (statistically significantly greater) in the presence of the candidate compound than in its absence, the candidate compound is identified as a stimulator of ORFX mRNA or protein expression. Alternatively, when expression of ORFX mRNA or protein is less (statistically significantly less) in the presence of the candidate

compound than in its absence, the candidate compound is identified as an inhibitor of ORFX mRNA or protein expression. The level of ORFX mRNA or protein expression in the cells can be determined by methods described herein for detecting ORFX mRNA or protein.

In yet another aspect of the invention, the ORFX proteins can be used as "bait proteins" in a two-hybrid assay or three hybrid assay (see, e.g., U.S. Pat. No. 5,283,317; Zervos et al. (1993) Cell 72:223-232; Madura et al. (1993) J Biol Chem 268:12046-12054; Bartel et al. (1993) Biotechniques 14:920-924; Iwabuchi et al. (1993) Oncogene 8:1693-1696; and Brent WO94/10300), to identify other proteins that bind to or interact with ORFX ("ORFX-binding proteins" or "ORFX-bp") and modulate ORFX activity. Such ORFX-binding proteins are also likely to be involved in the propagation of signals by the ORFX proteins as, for example, upstream or downstream elements of the ORFX pathway.

The two-hybrid system is based on the modular nature of most transcription factors, which consist of separable DNA-binding and activation domains. Briefly, the assay utilizes two different DNA constructs. In one construct, the gene that codes for ORFX is fused to a gene encoding the DNA binding domain of a known transcription factor (e.g., GAL-4). In the other construct, a DNA sequence, from a library of DNA sequences, that encodes an unidentified protein ("prey" or "sample") is fused to a gene that codes for the activation domain of the known transcription factor. If the "bait" and the "prey" proteins are able to interact, in vivo, forming a ORFX-dependent complex, the DNA-binding and activation domains of the transcription factor are brought into close proximity. This proximity allows transcription of a reporter gene (e.g., LacZ) that is operably linked to a transcriptional regulatory site responsive to the transcription factor. Expression of the reporter gene can be detected and cell colonies containing the functional transcription factor can be isolated and used to obtain the cloned gene that encodes the protein which interacts with ORFX.

This invention further pertains to novel agents identified by the above-described screening assays and uses thereof for treatments as described herein.

### **Detection Assays**

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Portions or fragments of the cDNA sequences identified herein (and the corresponding complete gene sequences) can be used in numerous ways as polynucleotide reagents. For example, these sequences can be used to: (i) map their respective genes on a chromosome; and, thus, locate gene regions associated with genetic disease; (ii) identify an individual from a

minute biological sample (tissue typing); and (iii) aid in forensic identification of a biological sample.

The ORFX sequences of the present invention can also be used to identify individuals from minute biological samples. In this technique, an individual's genomic DNA is digested with one or more restriction enzymes, and probed on a Southern blot to yield unique bands for identification. The sequences of the present invention are useful as additional DNA markers for RFLP ("restriction fragment length polymorphisms," described in U.S. Pat. No. 5,272,057).

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Furthermore, the sequences of the present invention can be used to provide an alternative technique that determines the actual base-by-base DNA sequence of selected portions of an individual's genome. Thus, the ORFX sequences described herein can be used to prepare two PCR primers from the 5' and 3' ends of the sequences. These primers can then be used to amplify an individual's DNA and subsequently sequence it.

Panels of corresponding DNA sequences from individuals, prepared in this manner, can provide unique individual identifications, as each individual will have a unique set of such DNA sequences due to allelic differences. The sequences of the present invention can be used to obtain such identification sequences from individuals and from tissue. The ORFX sequences of the invention uniquely represent portions of the human genome. Allelic variation occurs to some degree in the coding regions of these sequences, and to a greater degree in the noncoding regions. It is estimated that allelic variation between individual humans occurs with a frequency of about once per each 500 bases. Much of the allelic variation is due to single nucleotide polymorphisms (SNPs), which include restriction fragment length polymorphisms (RFLPs).

Each of the sequences described herein can, to some degree, be used as a standard against which DNA from an individual can be compared for identification purposes. Because greater numbers of polymorphisms occur in the noncoding regions, fewer sequences are necessary to differentiate individuals. The noncoding sequences of SEQ ID NO:2n-1 (wherein n=1 to 3161), as described above, can comfortably provide positive individual identification with a panel of perhaps 10 to 1,000 primers that each yield a noncoding amplified sequence of 100 bases. If predicted coding sequences are used, a more appropriate number of primers for positive individual identification would be 500-2,000.

### Predictive Medicine

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The present invention also pertains to the field of predictive medicine in which diagnostic assays, prognostic assays, pharmacogenomics, and monitoring clinical trials are used for prognostic (predictive) purposes to thereby treat an individual prophylactically. Accordingly, one aspect of the present invention relates to diagnostic assays for determining ORFX protein and/or nucleic acid expression as well as ORFX activity, in the context of a biological sample (e.g., blood, serum, cells, tissue) to thereby determine whether an individual is afflicted with a disease or disorder, or is at risk of developing a disorder, associated with aberrant ORFX expression or activity. The invention also provides for prognostic (or predictive) assays for determining whether an individual is at risk of developing a disorder associated with ORFX protein, nucleic acid expression or activity. For example, mutations in a ORFX gene can be assayed in a biological sample. Such assays can be used for prognostic or predictive purpose to thereby prophylactically treat an individual prior to the onset of a disorder characterized by or associated with ORFX protein, nucleic acid expression or activity.

Another aspect of the invention provides methods for determining ORFX protein, nucleic acid expression or ORFX activity in an individual to thereby select appropriate therapeutic or prophylactic agents for that individual (referred to herein as "pharmacogenomics"). Pharmacogenomics allows for the selection of agents (e.g., drugs) for therapeutic or prophylactic treatment of an individual based on the genotype of the individual (e.g., the genotype of the individual examined to determine the ability of the individual to respond to a particular agent.)

Yet another aspect of the invention pertains to monitoring the influence of agents (e.g., drugs, compounds) on the expression or activity of ORFX in clinical trials.

# Use of Partial ORFX Sequences in Forensic Biology

DNA-based identification techniques can also be used in forensic biology. Forensic biology is a scientific field employing genetic typing of biological evidence found at a crime scene as a means for positively identifying, for example, a perpetrator of a crime. To make such an identification, PCR technology can be used to amplify DNA sequences taken from very small biological samples such as tissues, e.g., hair or skin, or body fluids, e.g., blood, saliva, or semen

found at a crime scene. The amplified sequence can then be compared to a standard, thereby allowing identification of the origin of the biological sample.

The sequences of the present invention can be used to provide polynucleotide reagents, e.g., PCR primers, targeted to specific loci in the human genome, that can enhance the reliability of DNA-based forensic identifications by, for example, providing another "identification marker" (i.e. another DNA sequence that is unique to a particular individual). As mentioned above, actual base sequence information can be used for identification as an accurate alternative to patterns formed by restriction enzyme generated fragments. Sequences targeted to noncoding regions of SEQ ID NOs: \_\_ are particularly appropriate for this use as greater numbers of polymorphisms occur in the noncoding regions, making it easier to differentiate individuals using this technique. Examples of polynucleotide reagents include the ORFX sequences or portions thereof, e.g., fragments derived from the noncoding regions of one or more of SEQ ID NO:2n-1 (where n = 1 to 3161), having a length of at least 20 bases, preferably at least 30 bases.

The ORFX sequences described herein can further be used to provide polynucleotide reagents, e.g., labeled or label-able probes that can be used, for example, in an in situ hybridization technique, to identify a specific tissue, e.g., brain tissue, etc. This can be very useful in cases where a forensic pathologist is presented with a tissue of unknown origin. Panels of such ORFX probes can be used to identify tissue by species and/or by organ type.

In a similar fashion, these reagents, e.g., ORFX primers or probes can be used to screen tissue culture for contamination (i.e. screen for the presence of a mixture of different types of cells in a culture).

### **Predictive Medicine**

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The present invention also pertains to the field of predictive medicine in which diagnostic assays, prognostic assays, pharmacogenomics, and monitoring clinical trials are used for prognostic (predictive) purposes to thereby treat an individual prophylactically. Accordingly, one aspect of the present invention relates to diagnostic assays for determining ORFX protein and/or nucleic acid expression as well as ORFX activity, in the context of a biological sample (e.g., blood, serum, cells, tissue) to thereby determine whether an individual is afflicted with a disease or disorder, or is at risk of developing a disorder, associated with aberrant ORFX expression or activity. The invention also provides for prognostic (or predictive) assays for determining whether an individual is at risk of developing a disorder associated with ORFX protein, nucleic

acid expression or activity. For example, mutations in a ORFX gene can be assayed in a biological sample. Such assays can be used for prognostic or predictive purpose to thereby prophylactically treat an individual prior to the onset of a disorder characterized by or associated with ORFX protein, nucleic acid expression or activity.

Another aspect of the invention provides methods for determining ORFX protein, nucleic acid expression or ORFX activity in an individual to thereby select appropriate therapeutic or prophylactic agents for that individual (referred to herein as "pharmacogenomics"). Pharmacogenomics allows for the selection of agents (e.g., drugs) for therapeutic or prophylactic treatment of an individual based on the genotype of the individual (e.g., the genotype of the individual examined to determine the ability of the individual to respond to a particular agent.)

Yet another aspect of the invention pertains to monitoring the influence of agents (e.g., drugs, compounds) on the expression or activity of ORFX in clinical trials.

These and other agents are described in further detail in the following sections.

# **Diagnostic Assays**

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Other conditions in which proliferation of cells plays a role include tumors, restenosis, psoriasis, Dupuytren's contracture, diabetic complications, Kaposi's sarcoma and rheumatoid arthritis.

An ORFX polypeptide may be used to identify an interacting polypeptide a sample or tissue. The method comprises contacting the sample or tissue with ORFX, allowing formation of a complex between the ORFX polypeptide and the interacting polypeptide, and detecting the complex, if present.

The proteins of the invention may be used to stimulate production of antibodies specifically binding the proteins. Such antibodies may be used in immunodiagnostic procedures to detect the occurrence of the protein in a sample. The proteins of the invention may be used to stimulate cell growth and cell proliferation in conditions in which such growth would be favorable. An example would be to counteract toxic side effects of chemotherapeutic agents on, for example, hematopoiesis and platelet formation, linings of the gastrointestinal tract, and hair follicles. They may also be used to stimulate new cell growth in neurological disorders including, for example, Alzheimer's disease. Alternatively, antagonistic treatments may be administered in which an antibody specifically binding the ORFX -like proteins of the invention

would abrogate the specific growth-inducing effects of the proteins. Such antibodies may be useful, for example, in the treatment of proliferative disorders including various tumors and benign hyperplasias.

Polynucleotides or oligonucleotides corresponding to any one portion of the ORFX nucleic acids of SEQ ID NO:2n-1 (wherein n=1 to 3161) may be used to detect DNA containing a corresponding ORF gene, or detect the expression of a corresponding ORFX gene, or ORFX-like gene. For example, an ORFX nucleic acid expressed in a particular cell or tissue, as noted in Table 2, can be used to identify the presence of that particular cell type.

An exemplary method for detecting the presence or absence of ORFX in a biological sample involves obtaining a biological sample from a test subject and contacting the biological sample with a compound or an agent capable of detecting ORFX protein or nucleic acid (e.g., mRNA, genomic DNA) that encodes ORFX protein such that the presence of ORFX is detected in the biological sample. An agent for detecting ORFX mRNA or genomic DNA is a labeled nucleic acid probe capable of hybridizing to ORFX mRNA or genomic DNA. The nucleic acid probe can be, for example, a full-length ORFX nucleic acid, such as the nucleic acid of SEQ ID NO:2n-1 (wherein n = 1 to 3161), or a portion thereof, such as an oligonucleotide of at least 15, 30, 50, 100, 250 or 500 nucleotides in length and sufficient to specifically hybridize under stringent conditions to ORFX mRNA or genomic DNA, as described above. Other suitable probes for use in the diagnostic assays of the invention are described herein.

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An agent for detecting ORFX protein is an antibody capable of binding to ORFX protein, preferably an antibody with a detectable label. Antibodies can be polyclonal, or more preferably, monoclonal. An intact antibody, or a fragment thereof (e.g., Fab or F(ab')<sub>2</sub>) can be used. The term "labeled", with regard to the probe or antibody, is intended to encompass direct labeling of the probe or antibody by coupling (i.e., physically linking) a detectable substance to the probe or antibody, as well as indirect labeling of the probe or antibody by reactivity with another reagent that is directly labeled. Examples of indirect labeling include detection of a primary antibody using a fluorescently labeled secondary antibody and end-labeling of a DNA probe with biotin such that it can be detected with fluorescently labeled streptavidin. The term "biological sample" is intended to include tissues, cells and biological fluids isolated from a subject, as well as tissues, cells and fluids present within a subject. That is, the detection method of the invention can be used to detect ORFX mRNA, protein, or genomic DNA in a biological sample in vitro as

well as in vivo. For example, in vitro techniques for detection of ORFX mRNA include Northern hybridizations and in situ hybridizations. In vitro techniques for detection of ORFX protein include enzyme linked immunosorbent assays (ELISAs), Western blots, immunoprecipitations and immunofluorescence. In vitro techniques for detection of ORFX genomic DNA include Southern hybridizations. Furthermore, in vivo techniques for detection of ORFX protein include introducing into a subject a labeled anti-ORFX antibody. For example, the antibody can be labeled with a radioactive marker whose presence and location in a subject can be detected by standard imaging techniques.

In one embodiment, the biological sample contains protein molecules from the test subject. Alternatively, the biological sample can contain mRNA molecules from the test subject or genomic DNA molecules from the test subject. A preferred biological sample is a peripheral blood leukocyte sample isolated by conventional means from a subject.

In another embodiment, the methods further involve obtaining a control biological sample from a control subject, contacting the control sample with a compound or agent capable of detecting ORFX protein, mRNA, or genomic DNA, such that the presence of ORFX protein, mRNA or genomic DNA is detected in the biological sample, and comparing the presence of ORFX protein, mRNA or genomic DNA in the control sample with the presence of ORFX protein, mRNA or genomic DNA in the test sample.

The invention also encompasses kits for detecting the presence of ORFX in a biological sample. For example, the kit can comprise: a labeled compound or agent capable of detecting ORFX protein or mRNA in a biological sample; means for determining the amount of ORFX in the sample; and means for comparing the amount of ORFX in the sample with a standard. The compound or agent can be packaged in a suitable container. The kit can further comprise instructions for using the kit to detect ORFX protein or nucleic acid.

## **Prognostic Assays**

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The diagnostic methods described herein can furthermore be utilized to identify subjects having or at risk of developing a disease or disorder associated with aberrant ORFX expression or activity. For example, the assays described herein, such as the preceding diagnostic assays or the following assays, can be utilized to identify a subject having or at risk of developing a disorder associated with ORFX protein, nucleic acid expression or activity in, e.g., proliferative or differentiative disorders such as hyperplasias, tumors, restenosis, psoriasis, Dupuytren's

contracture, diabetic complications, or rheumatoid arthritis, etc.; and glia-associated disorders such as cerebral lesions, diabetic neuropathies, cerebral edema, senile dementia, Alzheimer's disease, etc. Alternatively, the prognostic assays can be utilized to identify a subject having or at risk for developing a disease or disorder. Thus, the present invention provides a method for identifying a disease or disorder associated with aberrant ORFX expression or activity in which a test sample is obtained from a subject and ORFX protein or nucleic acid (e.g., mRNA, genomic DNA) is detected, wherein the presence of ORFX protein or nucleic acid is diagnostic for a subject having or at risk of developing a disease or disorder associated with aberrant ORFX expression or activity. As used herein, a "test sample" refers to a biological sample obtained from a subject of interest. For example, a test sample can be a biological fluid (e.g., serum), cell sample, or tissue.

Furthermore, the prognostic assays described herein can be used to determine whether a subject can be administered an agent (e.g., an agonist, antagonist, peptidomimetic, protein, peptide, nucleic acid, small molecule, or other drug candidate) to treat a disease or disorder associated with aberrant ORFX expression or activity. For example, such methods can be used to determine whether a subject can be effectively treated with an agent for a disorder, such as a proliferative disorder, differentiative disorder, glia-associated disorders, etc. Thus, the present invention provides methods for determining whether a subject can be effectively treated with an agent for a disorder associated with aberrant ORFX expression or activity in which a test sample is obtained and ORFX protein or nucleic acid is detected (e.g., wherein the presence of ORFX protein or nucleic acid is diagnostic for a subject that can be administered the agent to treat a disorder associated with aberrant ORFX expression or activity.)

The methods of the invention can also be used to detect genetic lesions in a ORFX gene, thereby determining if a subject with the lesioned gene is at risk for, or suffers from, a proliferative disorder, differentiative disorder, glia-associated disorder, etc. In various embodiments, the methods include detecting, in a sample of cells from the subject, the presence or absence of a genetic lesion characterized by at least one of an alteration affecting the integrity of a gene encoding a ORFX-protein, or the mis-expression of the ORFX gene. For example, such genetic lesions can be detected by ascertaining the existence of at least one of (1) a deletion of one or more nucleotides from a ORFX gene; (2) an addition of one or more nucleotides to a ORFX gene; (3) a substitution of one or more nucleotides of a ORFX gene, (4) a chromosomal

rearrangement of a ORFX gene; (5) an alteration in the level of a messenger RNA transcript of a ORFX gene, (6) aberrant modification of a ORFX gene, such as of the methylation pattern of the genomic DNA, (7) the presence of a non-wild type splicing pattern of a messenger RNA transcript of a ORFX gene, (8) a non-wild type level of a ORFX-protein, (9) allelic loss of a ORFX gene, and (10) inappropriate post-translational modification of a ORFX-protein. As described herein, there are a large number of assay techniques known in the art which can be used for detecting lesions in a ORFX gene. A preferred biological sample is a peripheral blood leukocyte sample isolated by conventional means from a subject. However, any biological sample containing nucleated cells may be used, including, for example, buccal mucosal cells.

In certain embodiments, detection of the lesion involves the use of a probe/primer in a polymerase chain reaction (PCR) (see, e.g., U.S. Pat. Nos. 4,683,195 and 4,683,202), such as anchor PCR or RACE PCR, or, alternatively, in a ligation chain reaction (LCR) (see, e.g., Landegran et al. (1988) Science 241:1077-1080; and Nakazawa et al. (1994) PNAS 91:360-364), the latter of which can be particularly useful for detecting point mutations in the ORFX-gene (see Abravaya et al. (1995) Nucl Acids Res 23:675-682). This method can include the steps of collecting a sample of cells from a patient, isolating nucleic acid (e.g., genomic, mRNA or both) from the cells of the sample, contacting the nucleic acid sample with one or more primers that specifically hybridize to a ORFX gene under conditions such that hybridization and amplification of the ORFX gene (if present) occurs, and detecting the presence or absence of an amplification product, or detecting the size of the amplification product and comparing the length to a control sample. It is anticipated that PCR and/or LCR may be desirable to use as a preliminary amplification step in conjunction with any of the techniques used for detecting mutations described herein.

Alternative amplification methods include: self sustained sequence replication (Guatelli et al., 1990, Proc Natl Acad Sci USA 87:1874-1878), transcriptional amplification system (Kwoh, et al., 1989, Proc Natl Acad Sci USA 86:1173-1177), Q-Beta Replicase (Lizardi et al., 1988, BioTechnology 6:1197), or any other nucleic acid amplification method, followed by the detection of the amplified molecules using techniques well known to those of skill in the art. These detection schemes are especially useful for the detection of nucleic acid molecules if such molecules are present in very low numbers.

In an alternative embodiment, mutations in a ORFX gene from a sample cell can be identified by alterations in restriction enzyme cleavage patterns. For example, sample and control DNA is isolated, amplified (optionally), digested with one or more restriction endonucleases, and fragment length sizes are determined by gel electrophoresis and compared. Differences in fragment length sizes between sample and control DNA indicates mutations in the sample DNA. Moreover, the use of sequence specific ribozymes (see, for example, U.S. Pat. No. 5,493,531) can be used to score for the presence of specific mutations by development or loss of a ribozyme cleavage site.

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In other embodiments, genetic mutations in ORFX can be identified by hybridizing a sample and control nucleic acids, e.g., DNA or RNA, to high density arrays containing hundreds or thousands of oligonucleotides probes (Cronin et al. (1996) Human Mutation 7: 244-255; Kozal et al. (1996) Nature Medicine 2: 753-759). For example, genetic mutations in ORFX can be identified in two dimensional arrays containing light-generated DNA probes as described in Cronin et al. above. Briefly, a first hybridization array of probes can be used to scan through long stretches of DNA in a sample and control to identify base changes between the sequences by making linear arrays of sequential overlapping probes. This step allows the identification of point mutations. This step is followed by a second hybridization array that allows the characterization of specific mutations by using smaller, specialized probe arrays complementary to all variants or mutations detected. Each mutation array is composed of parallel probe sets, one complementary to the wild-type gene and the other complementary to the mutant gene.

In yet another embodiment, any of a variety of sequencing reactions known in the art can be used to directly sequence the ORFX gene and detect mutations by comparing the sequence of the sample ORFX with the corresponding wild-type (control) sequence. Examples of sequencing reactions include those based on techniques developed by Maxim and Gilbert (1977) *PNAS* 74:560 or Sanger (1977) *PNAS* 74:5463. It is also contemplated that any of a variety of automated sequencing procedures can be utilized when performing the diagnostic assays (Naeve et al., (1995) Biotechniques 19:448), including sequencing by mass spectrometry (see, e.g., PCT International Publ. No. WO 94/16101; Cohen et al. (1996) Adv Chromatogr 36:127-162; and Griffin et al. (1993) Appl Biochem Biotechnol 38:147-159).

Other methods for detecting mutations in the ORFX gene include methods in which protection from cleavage agents is used to detect mismatched bases in RNA/RNA or RNA/DNA

heteroduplexes (Myers et al. (1985) Science 230:1242). In general, the art technique of "mismatch cleavage" starts by providing heteroduplexes of formed by hybridizing (labeled) RNA or DNA containing the wild-type ORFX sequence with potentially mutant RNA or DNA obtained from a tissue sample. The double-stranded duplexes are treated with an agent that cleaves single-stranded regions of the duplex such as which will exist due to basepair mismatches between the control and sample strands. For instance, RNA/DNA duplexes can be treated with RNase and DNA/DNA hybrids treated with S1 nuclease to enzymatically digesting the mismatched regions. In other embodiments, either DNA/DNA or RNA/DNA duplexes can be treated with hydroxylamine or osmium tetroxide and with piperidine in order to digest mismatched regions. After digestion of the mismatched regions, the resulting material is then separated by size on denaturing polyacrylamide gels to determine the site of mutation. See, for example, Cotton et al (1988) Proc Natl Acad Sci USA 85:4397; Saleeba et al (1992) Methods Enzymol 217:286-295. In an embodiment, the control DNA or RNA can be labeled for detection.

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In still another embodiment, the mismatch cleavage reaction employs one or more proteins that recognize mismatched base pairs in double-stranded DNA (so called "DNA mismatch repair" enzymes) in defined systems for detecting and mapping point mutations in ORFX cDNAs obtained from samples of cells. For example, the mutY enzyme of *E. coli* cleaves A at G/A mismatches and the thymidine DNA glycosylase from HeLa cells cleaves T at G/T mismatches (Hsu et al. (1994) Carcinogenesis 15:1657-1662). According to an exemplary embodiment, a probe based on a ORFX sequence, e.g., a wild-type ORFX sequence, is hybridized to a cDNA or other DNA product from a test cell(s). The duplex is treated with a DNA mismatch repair enzyme, and the cleavage products, if any, can be detected from electrophoresis protocols or the like. See, for example, U.S. Pat. No. 5,459,039.

In other embodiments, alterations in electrophoretic mobility will be used to identify mutations in ORFX genes. For example, single strand conformation polymorphism (SSCP) may be used to detect differences in electrophoretic mobility between mutant and wild type nucleic acids (Orita et al. (1989) Proc Natl Acad Sci USA: 86:2766, see also Cotton (1993) Mutat Res 285:125-144; Hayashi (1992) Genet Anal Tech Appl 9:73-79). Single-stranded DNA fragments of sample and control ORFX nucleic acids will be denatured and allowed to renature. The secondary structure of single-stranded nucleic acids varies according to sequence, the resulting

alteration in electrophoretic mobility enables the detection of even a single base change. The DNA fragments may be labeled or detected with labeled probes. The sensitivity of the assay may be enhanced by using RNA, rather than DNA, in which the secondary structure is more sensitive to a change in sequence. In one embodiment, the subject method utilizes heteroduplex analysis to separate double stranded heteroduplex molecules on the basis of changes in electrophoretic mobility. See, e.g., Keen et al. (1991) Trends Genet 7:5.

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In yet another embodiment the movement of mutant or wild-type fragments in polyacrylamide gels containing a gradient of denaturant is assayed using denaturing gradient gel electrophoresis (DGGE). See, e.g., Myers et al (1985) Nature 313:495. When DGGE is used as the method of analysis, DNA will be modified to insure that it does not completely denature, for example by adding a GC clamp of approximately 40 bp of high-melting GC-rich DNA by PCR. In a further embodiment, a temperature gradient is used in place of a denaturing gradient to identify differences in the mobility of control and sample DNA. See, e.g., Rosenbaum and Reissner (1987) Biophys Chem 265:12753.

Examples of other techniques for detecting point mutations include, but are not limited to, selective oligonucleotide hybridization, selective amplification, or selective primer extension. For example, oligonucleotide primers may be prepared in which the known mutation is placed centrally and then hybridized to target DNA under conditions that permit hybridization only if a perfect match is found. See, e.g., Saiki et al. (1986) Nature 324:163); Saiki et al. (1989) Proc Natl Acad. Sci USA 86:6230. Such allele specific oligonucleotides are hybridized to PCR amplified target DNA or a number of different mutations when the oligonucleotides are attached to the hybridizing membrane and hybridized with labeled target DNA.

Alternatively, allele specific amplification technology that depends on selective PCR amplification may be used in conjunction with the instant invention. Oligonucleotides used as primers for specific amplification may carry the mutation of interest in the center of the molecule (so that amplification depends on differential hybridization) (Gibbs et al. (1989) Nucleic Acids Res 17:2437-2448) or at the extreme 3' end of one primer where, under appropriate conditions, mismatch can prevent, or reduce polymerase extension (Prossner (1993) Tibtech 11:238). In addition it may be desirable to introduce a novel restriction site in the region of the mutation to create cleavage-based detection. See, e.g., Gasparini et al (1992) Mol Cell Probes 6:1. It is anticipated that in certain embodiments amplification may also be performed using Taq ligase

for amplification. See, e.g., Barany (1991) Proc Natl Acad Sci USA 88:189. In such cases, ligation will occur only if there is a perfect match at the 3' end of the 5' sequence, making it possible to detect the presence of a known mutation at a specific site by looking for the presence or absence of amplification.

The methods described herein may be performed, for example, by utilizing pre-packaged diagnostic kits comprising at least one probe nucleic acid or antibody reagent described herein, which may be conveniently used, e.g., in clinical settings to diagnose patients exhibiting symptoms or family history of a disease or illness involving a ORFX gene.

Furthermore, any cell type or tissue, preferably peripheral blood leukocytes, in which ORFX is expressed may be utilized in the prognostic assays described herein. However, any biological sample containing nucleated cells may be used, including, for example, buccal mucosal cells.

## **Pharmacogenomics**

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Agents, or modulators that have a stimulatory or inhibitory effect on ORFX activity (e.g., ORFX gene expression), as identified by a screening assay described herein can be administered to individuals to treat (prophylactically or therapeutically) disorders (e.g., neurological, cancerrelated or gestational disorders) associated with aberrant ORFX activity. In conjunction with such treatment, the pharmacogenomics (i.e., the study of the relationship between an individual's genotype and that individual's response to a foreign compound or drug) of the individual may be considered. Differences in metabolism of therapeutics can lead to severe toxicity or therapeutic failure by altering the relation between dose and blood concentration of the pharmacologically active drug. Thus, the pharmacogenomics of the individual permits the selection of effective agents (e.g., drugs) for prophylactic or therapeutic treatments based on a consideration of the individual's genotype. Such pharmacogenomics can further be used to determine appropriate dosages and therapeutic regimens. Accordingly, the activity of ORFX protein, expression of ORFX nucleic acid, or mutation content of ORFX genes in an individual can be determined to thereby select appropriate agent(s) for therapeutic or prophylactic treatment of the individual.

Pharmacogenomics deals with clinically significant hereditary variations in the response to drugs due to altered drug disposition and abnormal action in affected persons. See e.g., Eichelbaum, 1996, Clin Exp Pharmacol Physiol, 23:983-985 and Linder, 1997, Clin Chem, 43:254-266. In general, two types of pharmacogenetic conditions can be differentiated. Genetic

conditions transmitted as a single factor altering the way drugs act on the body (altered drug action) or genetic conditions transmitted as single factors altering the way the body acts on drugs (altered drug metabolism). These pharmacogenetic conditions can occur either as rare defects or as polymorphisms. For example, glucose-6-phosphate dehydrogenase (G6PD) deficiency is a common inherited enzymopathy in which the main clinical complication is haemolysis after ingestion of oxidant drugs (anti-malarials, sulfonamides, analgesics, nitrofurans) and consumption of fava beans.

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As an illustrative embodiment, the activity of drug metabolizing enzymes is a major determinant of both the intensity and duration of drug action. The discovery of genetic polymorphisms of drug metabolizing enzymes (e.g., N-acetyltransferase 2 (NAT 2) and cytochrome P450 enzymes CYP2D6 and CYP2C19) has provided an explanation as to why some patients do not obtain the expected drug effects or show exaggerated drug response and serious toxicity after taking the standard and safe dose of a drug. These polymorphisms are expressed in two phenotypes in the population, the extensive metabolizer (EM) and poor metabolizer (PM). The prevalence of PM is different among different populations. For example, the gene coding for CYP2D6 is highly polymorphic and several mutations have been identified in PM, which all lead to the absence of functional CYP2D6. Poor metabolizers of CYP2D6 and CYP2C19 quite frequently experience exaggerated drug response and side effects when they receive standard doses. If a metabolite is the active therapeutic moiety, PM show no therapeutic response, as demonstrated for the analgesic effect of codeine mediated by its CYP2D6-formed metabolite morphine. The other extreme are the so called ultra-rapid metabolizers who do not respond to standard doses. Recently, the molecular basis of ultra-rapid metabolism has been identified to be due to CYP2D6 gene amplification.

Thus, the activity of ORFX protein, expression of ORFX nucleic acid, or mutation content of ORFX genes in an individual can be determined to thereby select appropriate agent(s) for therapeutic or prophylactic treatment of the individual. In addition, pharmacogenetic studies can be used to apply genotyping of polymorphic alleles encoding drug-metabolizing enzymes to the identification of an individual's drug responsiveness phenotype. This knowledge, when applied to dosing or drug selection, can avoid adverse reactions or therapeutic failure and thus enhance therapeutic or prophylactic efficiency when treating a subject with a ORFX modulator, such as a modulator identified by one of the exemplary screening assays described herein.

# Monitoring Clinical Efficacy

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Monitoring the influence of agents (e.g., drugs, compounds) on the expression or activity of ORFX (e.g., the ability to modulate aberrant cell proliferation and/or differentiation) can be applied in basic drug screening and in clinical trials. For example, the effectiveness of an agent determined by a screening assay as described herein to increase ORFX gene expression, protein levels, or upregulate ORFX activity, can be monitored in clinical trials of subjects exhibiting decreased ORFX gene expression, protein levels, or downregulated ORFX activity.

Alternatively, the effectiveness of an agent determined by a screening assay to decrease ORFX gene expression, protein levels, or downregulate ORFX activity, can be monitored in clinical trials of subjects exhibiting increased ORFX gene expression, protein levels, or upregulated ORFX activity. In such clinical trials, the expression or activity of ORFX and, preferably, other genes that have been implicated in, for example, a proliferative or neurological disorder, can be used as a "read out" or marker of the responsiveness of a particular cell.

For example, genes, including ORFX, that are modulated in cells by treatment with an agent (e.g., compound, drug or small molecule) that modulates ORFX activity (e.g., identified in a screening assay as described herein) can be identified. Thus, to study the effect of agents on cellular proliferation disorders, for example, in a clinical trial, cells can be isolated and RNA prepared and analyzed for the levels of expression of ORFX and other genes implicated in the disorder. The levels of gene expression (i.e., a gene expression pattern) can be quantified by Northern blot analysis or RT-PCR, as described herein, or alternatively by measuring the amount of protein produced, by one of the methods as described herein, or by measuring the levels of activity of ORFX or other genes. In this way, the gene expression pattern can serve as a marker, indicative of the physiological response of the cells to the agent. Accordingly, this response state may be determined before, and at various points during, treatment of the individual with the agent.

In one embodiment, the invention provides a method for monitoring the effectiveness of treatment of a subject with an agent (e.g., an agonist, antagonist, protein, peptide, nucleic acid, peptidomimetic, small molecule, or other drug candidate identified by the screening assays described herein) comprising the steps of (i) obtaining a pre-administration sample from a subject prior to administration of the agent; (ii) detecting the level of expression of a ORFX protein, mRNA, or genomic DNA in the preadministration sample; (iii) obtaining one or more

post-administration samples from the subject; (iv) detecting the level of expression or activity of the ORFX protein, mRNA, or genomic DNA in the post-administration samples; (v) comparing the level of expression or activity of the ORFX protein, mRNA, or genomic DNA in the pre-administration sample with the ORFX protein, mRNA, or genomic DNA in the post administration sample or samples; and (vi) altering the administration of the agent to the subject accordingly. For example, increased administration of the agent may be desirable to increase the expression or activity of ORFX to higher levels than detected, i.e., to increase the effectiveness of the agent. Alternatively, decreased administration of the agent may be desirable to decrease expression or activity of ORFX to lower levels than detected, i.e., to decrease the effectiveness of the agent.

### Methods of Treatment

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The present invention provides for both prophylactic and therapeutic methods of treating a subject at risk of (or susceptible to) a disorder or having a disorder associated with aberrant ORFX expression or activity.

Diseases and disorders that are characterized by increased (relative to a subject not suffering from the disease or disorder) levels or biological activity may be treated with Therapeutics that antagonize (i.e., reduce or inhibit) activity. Therapeutics that antagonize activity may be administered in a therapeutic or prophylactic manner. Therapeutics that may be utilized include, but are not limited to, (i) a ORFX polypeptide, or analogs, derivatives, fragments or homologs thereof; (ii) antibodies to a ORFX peptide; (iii) nucleic acids encoding a ORFX peptide; (iv) administration of antisense nucleic acid and nucleic acids that are "dysfunctional" (i.e., due to a heterologous insertion within the coding sequences of coding sequences to a ORFX peptide) that are utilized to "knockout" endogenous function of a ORFX peptide by homologous recombination (see, e.g., Capecchi, 1989, Science 244: 1288-1292); or (v) modulators (i.e., inhibitors, agonists and antagonists, including additional peptide mimetic of the invention or antibodies specific to a peptide of the invention) that alter the interaction between a ORFX peptide and its binding partner.

Diseases and disorders that are characterized by decreased (relative to a subject not suffering from the disease or disorder) levels or biological activity may be treated with Therapeutics that increase (i.e., are agonists to) activity. Therapeutics that upregulate activity may be administered in a therapeutic or prophylactic manner. Therapeutics that may be utilized

include, but are not limited to, a ORFX peptide, or analogs, derivatives, fragments or homologs thereof; or an agonist that increases bioavailability.

Increased or decreased levels can be readily detected by quantifying peptide and/or RNA, by obtaining a patient tissue sample (e.g., from biopsy tissue) and assaying it in vitro for RNA or peptide levels, structure and/or activity of the expressed peptides (or mRNAs of a ORFX peptide). Methods that are well-known within the art include, but are not limited to, immunoassays (e.g., by Western blot analysis, immunoprecipitation followed by sodium dodecyl sulfate (SDS) polyacrylamide gel electrophoresis, immunocytochemistry, etc.) and/or hybridization assays to detect expression of mRNAs (e.g., Northern assays, dot blots, in situ hybridization, etc.).

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In one aspect, the invention provides a method for preventing, in a subject, a disease or condition associated with an aberrant ORFX expression or activity, by administering to the subject an agent that modulates ORFX expression or at least one ORFX activity. Subjects at risk for a disease that is caused or contributed to by aberrant ORFX expression or activity can be identified by, for example, any or a combination of diagnostic or prognostic assays as described herein. Administration of a prophylactic agent can occur prior to the manifestation of symptoms characteristic of the ORFX aberrancy, such that a disease or disorder is prevented or, alternatively, delayed in its progression. Depending on the type of ORFX aberrancy, for example, a ORFX agonist or ORFX antagonist agent can be used for treating the subject. The appropriate agent can be determined based on screening assays described herein.

Another aspect of the invention pertains to methods of modulating ORFX expression or activity for therapeutic purposes. The modulatory method of the invention involves contacting a cell with an agent that modulates one or more of the activities of ORFX protein activity associated with the cell. An agent that modulates ORFX protein activity can be an agent as described herein, such as a nucleic acid or a protein, a naturally-occurring cognate ligand of a ORFX protein, a peptide, a ORFX peptidomimetic, or other small molecule. In one embodiment, the agent stimulates one or more ORFX protein activity. Examples of such stimulatory agents include active ORFX protein and a nucleic acid molecule encoding ORFX that has been introduced into the cell. In another embodiment, the agent inhibits one or more ORFX protein activity. Examples of such inhibitory agents include antisense ORFX nucleic acid molecules and anti-ORFX antibodies. These modulatory methods can be performed *in vitro* 

(e.g., by culturing the cell with the agent) or, alternatively, in vivo (e.g., by administering the agent to a subject). As such, the present invention provides methods of treating an individual afflicted with a disease or disorder characterized by aberrant expression or activity of a ORFX protein or nucleic acid molecule. In one embodiment, the method involves administering an agent (e.g., an agent identified by a screening assay described herein), or combination of agents that modulates (e.g., upregulates or downregulates) ORFX expression or activity. In another embodiment, the method involves administering a ORFX protein or nucleic acid molecule as therapy to compensate for reduced or aberrant ORFX expression or activity.

## Determination of the Biological Effect of a Therapeutic

In various embodiments of the present invention, suitable *in vitro* or *in vivo* assays are utilized to determine the effect of a specific Therapeutic and whether its administration is indicated for treatment of the affected tissue.

In various specific embodiments, in vitro assays may be performed with representative cells of the type(s) involved in the patient's disorder, to determine if a given Therapeutic exerts the desired effect upon the cell type(s). Compounds for use in therapy may be tested in suitable animal model systems including, but not limited to rats, mice, chicken, cows, monkeys, rabbits, and the like, prior to testing in human subjects. Similarly, for in vivo testing, any of the animal model system known in the art may be used prior to administration to human subjects.

#### Malignancies

Some ORFX polypeptides are expressed in cancerous cells (see, e.g., Tables 1 and 2). Accordingly, the corresponding ORF protein is involved in the regulation of cell proliferation. Accordingly, Therapeutics of the present invention may be useful in the therapeutic or prophylactic treatment of diseases or disorders that are associated with cell hyperproliferation and/or loss of control of cell proliferation (e.g., cancers, malignancies and tumors). For a review of such hyperproliferation disorders, see e.g., Fishman, et al., 1985. MEDICINE, 2nd ed., J.B. Lippincott Co., Philadelphia, PA.

Therapeutics of the present invention may be assayed by any method known within the art for efficacy in treating or preventing malignancies and related disorders. Such assays include,

but are not limited to, *in vitro* assays utilizing transformed cells or cells derived from the patient's tumor, as well as *in vivo* assays using animal models of cancer or malignancies. Potentially effective Therapeutics are those that, for example, inhibit the proliferation of tumor-derived or transformed cells in culture or cause a regression of tumors in animal models, in comparison to the controls.

In the practice of the present invention, once a malignancy or cancer has been shown to be amenable to treatment by modulating (i.e., inhibiting, antagonizing or agonizing) activity, that cancer or malignancy may subsequently be treated or prevented by the administration of a Therapeutic that serves to modulate protein function.

# Premalignant conditions

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The Therapeutics of the present invention that are effective in the therapeutic or prophylactic treatment of cancer or malignancies may also be administered for the treatment of pre-malignant conditions and/or to prevent the progression of a pre-malignancy to a neoplastic or malignant state. Such prophylactic or therapeutic use is indicated in conditions known or suspected of preceding progression to neoplasia or cancer, in particular, where non-neoplastic cell growth consisting of hyperplasia, metaplasia or, most particularly, dysplasia has occurred. For a review of such abnormal cell growth see *e.g.*, Robbins & Angell, 1976. BASIC PATHOLOGY, 2nd ed., W.B. Saunders Co., Philadelphia, PA.

Hyperplasia is a form of controlled cell proliferation involving an increase in cell number in a tissue or organ, without significant alteration in its structure or function. For example, it has been demonstrated that endometrial hyperplasia often precedes endometrial cancer. Metaplasia is a form of controlled cell growth in which one type of mature or fully differentiated cell substitutes for another type of mature cell. Metaplasia may occur in epithelial or connective tissue cells. Dysplasia is generally considered a precursor of cancer, and is found mainly in the epithelia. Dysplasia is the most disorderly form of non-neoplastic cell growth, and involves a loss in individual cell uniformity and in the architectural orientation of cells. Dysplasia characteristically occurs where there exists chronic irritation or inflammation, and is often found in the cervix, respiratory passages, oral cavity, and gall bladder.

Alternatively, or in addition to the presence of abnormal cell growth characterized as hyperplasia, metaplasia, or dysplasia, the presence of one or more characteristics of a transformed or malignant phenotype displayed either *in vivo* or *in vitro* within a cell sample derived from a patient, is indicative of the desirability of prophylactic/therapeutic administration of a Therapeutic that possesses the ability to modulate activity of An aforementioned protein. Characteristics of a transformed phenotype include, but are not limited to: (*i*) morphological changes; (*ii*) looser substratum attachment; (*iii*) loss of cell-to-cell contact inhibition; (*iv*) loss of anchorage dependence; (*v*) protease release; (*vi*) increased sugar transport; (*vii*) decreased serum requirement; (*viii*) expression of fetal antigens, (*ix*) disappearance of the 250 kDal cell-surface protein, and the like. See *e.g.*, Richards, *et al.*, 1986. Molecular Pathology, W.B. Saunders Co., Philadelphia, PA.

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In a specific embodiment of the present invention, a patient that exhibits one or more of the following predisposing factors for malignancy is treated by administration of an effective amount of a Therapeutic: (i) a chromosomal translocation associated with a malignancy (e.g., the Philadelphia chromosome (bcr/abl) for chronic myelogenous leukemia and t(14;18) for follicular lymphoma, etc.); (ii) familial polyposis or Gardner's syndrome (possible forerunners of colon cancer); (iii) monoclonal gammopathy of undetermined significance (a possible precursor of multiple myeloma) and (iv) a first degree kinship with persons having a cancer or pre-cancerous disease showing a Mendelian (genetic) inheritance pattern (e.g., familial polyposis of the colon, Gardner's syndrome, hereditary exostosis, polyendocrine adenomatosis, Peutz-Jeghers syndrome, neurofibromatosis of Von Recklinghausen, medullary thyroid carcinoma with amyloid production and pheochromocytoma, retinoblastoma, carotid body tumor, cutaneous melanocarcinoma, intraocular melanocarcinoma, xeroderma pigmentosum, ataxia telangiectasia, Chediak-Higashi syndrome, albinism, Fanconi's aplastic anemia and Bloom's syndrome).

In another embodiment, a Therapeutic of the present invention is administered to a human patient to prevent the progression to breast, colon, lung, pancreatic, or uterine cancer, or melanoma or sarcoma.

# Hyperproliferative and dysproliferative disorders

In one embodiment of the present invention, a Therapeutic is administered in the therapeutic or prophylactic treatment of hyperproliferative or benign dysproliferative disorders. The efficacy in treating or preventing hyperproliferative diseases or disorders of a Therapeutic of the present invention may be assayed by any method known within the art. Such assays include in vitro cell proliferation assays, in vitro or in vivo assays using animal models of hyperproliferative diseases or disorders, or the like. Potentially effective Therapeutics may, for example, promote cell proliferation in culture or cause growth or cell proliferation in animal models in comparison to controls.

Specific embodiments of the present invention are directed to the treatment or prevention of cirrhosis of the liver (a condition in which scarring has overtaken normal liver regeneration processes); treatment of keloid (hypertrophic scar) formation causing disfiguring of the skin in which the scarring process interferes with normal renewal; psoriasis (a common skin condition characterized by excessive proliferation of the skin and delay in proper cell fate determination); benign tumors; fibrocystic conditions and tissue hypertrophy (e.g., benign prostatic hypertrophy).

# Neurodegenerative disorders

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Some ORFX proteins are found in cell types have been implicated in the deregulation of cellular maturation and apoptosis, which are both characteristic of neurodegenerative disease. Accordingly, Therapeutics of the invention, particularly but not limited to those that modulate (or supply) activity of an aforementioned protein, may be effective in treating or preventing neurodegenerative disease. Therapeutics of the present invention that modulate the activity of an aforementioned protein involved in neurodegenerative disorders can be assayed by any method known in the art for efficacy in treating or preventing such neurodegenerative diseases and disorders. Such assays include *in vitro* assays for regulated cell maturation or inhibition of apoptosis or *in vivo* assays using animal models of neurodegenerative diseases or disorders, or any of the assays described below. Potentially effective Therapeutics, for example but not by way of limitation, promote regulated cell maturation and prevent cell apoptosis in culture, or reduce neurodegeneration in animal models in comparison to controls.

Once a neurodegenerative disease or disorder has been shown to be amenable to treatment by modulation activity, that neurodegenerative disease or disorder can be treated or prevented by administration of a Therapeutic that modulates activity. Such diseases include all degenerative disorders involved with aging, especially osteoarthritis and neurodegenerative disorders.

### Disorders related to organ transplantation

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Some ORFX can be associated with disorders related to organ transplantation, in particular but not limited to organ rejection. Therapeutics of the invention, particularly those that modulate (or supply) activity, may be effective in treating or preventing diseases or disorders related to organ transplantation. Therapeutics of the invention (particularly Therapeutics that modulate the levels or activity of an aforementioned protein) can be assayed by any method known in the art for efficacy in treating or preventing such diseases and disorders related to organ transplantation. Such assays include *in vitro* assays for using cell culture models as described below, or *in vivo* assays using animal models of diseases and disorders related to organ transplantation, see *e.g.*, below. Potentially effective Therapeutics, for example but not by way of limitation, reduce immune rejection responses in animal models in comparison to controls.

Accordingly, once diseases and disorders related to organ transplantation are shown to be amenable to treatment by modulation of activity, such diseases or disorders can be treated or prevented by administration of a Therapeutic that modulates activity.

#### Cardiovascular Disease

GENX has been implicated in cardiovascular disorders, including in atherosclerotic plaque formation. Diseases such as cardiovascular disease, including cerebral thrombosis or hemorrhage, ischemic heart or renal disease, peripheral vascular disease, or thrombosis of other major vessel, and other diseases, including diabetes mellitus, hypertension, hypothyroidism, cholesterol ester storage disease, systemic lupus erythematosus, homocysteinemia, and familial protein or lipid processing diseases, and the like, are either directly or indirectly associated with atherosclerosis. Accordingly, Therapeutics of the invention, particularly those that modulate (or supply) activity or formation may be effective in treating or preventing

atherosclerosis-associated diseases or disorders. Therapeutics of the invention (particularly Therapeutics that modulate the levels or activity) can be assayed by any method known in the art, including those described below, for efficacy in treating or preventing such diseases and disorders.

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A vast array of animal and cell culture models exist for processes involved in atherosclerosis. A limited and non-exclusive list of animal models includes knockout mice for premature atherosclerosis (Kurabayashi and Yazaki, 1996, Int. Angiol. 15: 187-194), transgenic mouse models of atherosclerosis (Kappel et al., 1994, FASEB J. 8: 583-592), antisense oligonucleotide treatment of animal models (Callow, 1995, Curr. Opin. Cardiol. 10: 569-576), transgenic rabbit models for atherosclerosis (Taylor, 1997, Ann. N.Y. Acad. Sci 811: 146-152), hypercholesterolemic animal models (Rosenfeld, 1996, Diabetes Res. Clin. Pract. 30 Suppl.: 1-11), hyperlipidemic mice (Paigen et al., 1994, Curt. Opin. Lipidol. 5: 258-264), and inhibition of lipoxygenase in animals (Sigal et al., 1994, Ann. N.Y. Acad. Sci. 714: 211-224). In addition, in vitro cell models include but are not limited to monocytes exposed to low density lipoprotein (Frostegard et al., 1996, Atherosclerosis 121: 93-103), cloned vascular smooth muscle cells (Suttles et al., 1995, Exp. Cell Res. 218: 331-338), endothelial cell-derived chemoattractant exposed T cells (Katz et al., 1994, J. Leukoc. Biol. 55: 567-573), cultured human aortic endothelial cells (Farber et al., 1992, Am. J. Physiol. 262: H1088-1085), and foam cell cultures (Libby et al., 1996, Curr Opin Lipidol 7: 330-335). Potentially effective Therapeutics, for example but not by way of limitation, reduce foam cell formation in cell culture models, or reduce atherosclerotic plaque formation in hypercholesterolemic mouse models of atherosclerosis in comparison to controls.

Accordingly, once an atherosclerosis-associated disease or disorder has been shown to be amenable to treatment by modulation of activity or formation, that disease or disorder can be treated or prevented by administration of a Therapeutic that modulates activity.

# Cytokine and Cell Proliferation/Differentiation Activity

A GENX protein of the present invention may exhibit cytokine, cell proliferation (either inducing or inhibiting) or cell differentiation (either inducing or inhibiting) activity or may induce production of other cytokines in certain cell populations. Many protein factors discovered

to date, including all known cytokines, have exhibited activity in one or more factor dependent cell proliferation assays, and hence the assays serve as a convenient confirmation of cytokine activity. The activity of a protein of the present invention is evidenced by any one of a number of routine factor dependent cell proliferation assays for cell lines including, without limitation, 32D, DA2, DA1G, T10, B9, B9/11, BaF3, MC9/G, M+ (preB M+), 2E8, RB5, DA1, 123, T1165, HT2, CTLL2, TF-1, Mo7e and CMK.

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The activity of a protein of the invention may, among other means, be measured by the following methods: Assays for T-cell or thymocyte proliferation include without limitation those described in: Current Protocols in Immunology, Ed by Coligan et al., Greene Publishing Associates and Wiley-Interscience (Chapter 3 and Chapter 7); Takai et al., J Immunol 137:3494-3500, 1986; Bertagnoili et al., J Immunol 145:1706-1712, 1990; Bertagnolli et al., Cell Immunol 133:327-341, 1991; Bertagnolli, et al., J Immunol 149:3778-3783, 1992; Bowman et al., J Immunol 152:1756-1761, 1994.

Assays for cytokine production and/or proliferation of spleen cells, lymph node cells or thymocytes include, without limitation, those described by Kruisbeek and Shevach, In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan *et al.*, eds. Vol 1, pp. 3.12.1-14, John Wiley and Sons, Toronto 1994; and by Schreiber, In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan eds. Vol 1 pp. 6.8.1-8, John Wiley and Sons, Toronto 1994.

Assays for proliferation and differentiation of hematopoietic and lymphopoietic cells include, without limitation, those described by Bottomly et al., In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan et al., eds. Vol 1 pp. 6.3.1-6.3.12, John Wiley and Sons, Toronto 1991; deVries et al., J Exp Med 173:1205-1211, 1991; Moreau et al., Nature 336:690-692, 1988; Greenberger et al., Proc Natl Acad Sci U.S.A. 80:2931-2938, 1983; Nordan, In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan et al., eds. Vol 1 pp. 6.6.1-5, John Wiley and Sons, Toronto 1991; Smith et al., Proc Natl Acad Sci U.S.A. 83:1857-1861, 1986; Measurement of human Interleukin 11-Bennett, et al. In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan et al., eds. Vol 1 pp. 6.15.1 John Wiley and Sons, Toronto 1991; Ciarletta, et al., In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan et al., eds. Vol 1 pp. 6.13.1, John Wiley and Sons, Toronto 1991.

Assays for T-cell clone responses to antigens (which will identify, among others, proteins that affect APC-T cell interactions as well as direct T-cell effects by measuring proliferation and

cytokine production) include, without limitation, those described In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan et al., eds., Greene Publishing Associates and Wiley-Interscience (Chapter 3Chapter 6, Chapter 7); Weinberger et al., Proc Natl Acad Sci USA 77:6091-6095, 1980; Weinberger et al., Eur J Immun 11:405-411, 1981; Takai et al., J Immunol 137:3494-3500, 1986; Takai et al., J Immunol 140:508-512, 1988.

# Immune Stimulating or Suppressing Activity

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A GENX protein of the present invention may also exhibit immune stimulating or immune suppressing activity, including without limitation the activities for which assays are described herein. A protein may be useful in the treatment of various immune deficiencies and disorders (including severe combined immunodeficiency (SCID)), e.g., in regulating (up or down) growth and proliferation of T and/or B lymphocytes, as well as effecting the cytolytic activity of NK cells and other cell populations. These immune deficiencies may be genetic or be caused by vital (e.g., HIV) as well as bacterial or fungal infections, or may result from autoimmune disorders. More specifically, infectious diseases causes by vital, bacterial, fungal or other infection may be treatable using a protein of the present invention, including infections by HIV, hepatitis viruses, herpesviruses, mycobacteria, Leishmania species., malaria species. and various fungal infections such as candidiasis. Of course, in this regard, a protein of the present invention may also be useful where a boost to the immune system generally may be desirable, i.e., in the treatment of cancer.

Autoimmune disorders which may be treated using a protein of the present invention include, for example, connective tissue disease, multiple sclerosis, systemic lupus erythematosus, rheumatoid arthritis, autoimmune pulmonary inflammation, Guillain-Barre syndrome, autoimmune thyroiditis, insulin dependent diabetes mellitus, myasthenia gravis, graft-versus-host disease and autoimmune inflammatory eye disease. Such a protein of the present invention may also to be useful in the treatment of allergic reactions and conditions, such as asthma (particularly allergic asthma) or other respiratory problems. Other conditions, in which immune suppression is desired (including, for example, organ transplantation), may also be treatable using a protein of the present invention.

Using the proteins of the invention it may also be possible to immune responses, in a number of ways. Down regulation may be in the form of inhibiting or blocking an immune response already in progress or may involve preventing the induction of an immune response.

The functions of activated T cells may be inhibited by suppressing T cell responses or by inducing specific tolerance in T cells, or both. Immunosuppression of T cell responses is generally an active, non-antigen-specific, process which requires continuous exposure of the T cells to the suppressive agent. Tolerance, which involves inducing non-responsiveness or energy in T cells, is distinguishable from immunosuppression in that it is generally antigen-specific and persists after exposure to the tolerizing agent has ceased. Operationally, tolerance can be demonstrated by the lack of a T cell response upon re-exposure to specific antigen in the absence of the tolerizing agent.

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Down regulating or preventing one or more antigen functions (including without limitation B lymphocyte antigen functions (such as, for example, B7), e.g., preventing high level lymphokine synthesis by activated T cells, will be useful in situations of tissue, skin and organ transplantation and in graft-versus-host disease (GVHD). For example, blockage of T cell function should result in reduced tissue destruction in tissue transplantation. Typically, in tissue transplants, rejection of the transplant is initiated through its recognition as foreign by T cells, followed by an immune reaction that destroys the transplant. The administration of a molecule which inhibits or blocks interaction of a B7 lymphocyte antigen with its natural ligand(s) on immune cells (such as a soluble, monomeric form of a peptide having B7-2 activity alone or in conjunction with a monomeric form of a peptide having an activity of another B lymphocyte antigen (e.g., B7-1, B7-3) or blocking antibody), prior to transplantation can lead to the binding of the molecule to the natural ligand(s) on the immune cells without transmitting the corresponding costimulatory signal. Blocking B lymphocyte antigen function in this matter prevents cytokine synthesis by immune cells, such as T cells, and thus acts as an immunosuppressant. Moreover, the lack of costimulation may also be sufficient to energize the T cells, thereby inducing tolerance in a subject. Induction of long-term tolerance by B lymphocyte antigen-blocking reagents may avoid the necessity of repeated administration of these blocking reagents. To achieve sufficient immunosuppression or tolerance in a subject, it may also be necessary to block the function of B lymphocyte antigens.

The efficacy of particular blocking reagents in preventing organ transplant rejection or GVHD can be assessed using animal models that are predictive of efficacy in humans. Examples of appropriate systems which can be used include allogeneic cardiac grafts in rats and xenogeneic pancreatic islet cell grafts in mice, both of which have been used to examine the

immunosuppressive effects of CTLA4Ig fusion proteins in vivo as described in Lenschow et al., Science 257:789-792 (1992) and Turka et al., Proc Natl Acad Sci USA, 89:11102-11105 (1992). In addition, murine models of GVHD (see Paul ed., Fundamental Immunology, Raven Press, New York, 1989, pp. 846-847) can be used to determine the effect of blocking B lymphocyte antigen function in vivo on the development of that disease.

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Blocking antigen function may also be therapeutically useful for treating autoimmune diseases. Many autoimmune disorders are the result of inappropriate activation of T cells that are reactive against self tissue and which promote the production of cytokines and auto-antibodies involved in the pathology of the diseases. Preventing the activation of autoreactive T cells may reduce or eliminate disease symptoms. Administration of reagents which block costimulation of T cells by disrupting receptor: ligand interactions of B lymphocyte antigens can be used to inhibit T cell activation and prevent production of auto-antibodies or T cell-derived cytokines which may be involved in the disease process. Additionally, blocking reagents may induce antigen-specific tolerance of autoreactive T cells which could lead to long-term relief from the disease. The efficacy of blocking reagents in preventing or alleviating autoimmune disorders can be determined using a number of well-characterized animal models of human autoimmune diseases. Examples include murine experimental autoimmune encephalitis, systemic lupus erythematosis in MRL/lpr/lpr mice or NZB hybrid mice, murine autoimmune collagen arthritis, diabetes mellitus in NOD mice and BB rats, and murine experimental myasthenia gravis (see Paul ed., Fundamental Immunology, Raven Press, New York, 1989, pp. 840-856).

Upregulation of an antigen function (preferably a B lymphocyte antigen function), as a means of up regulating immune responses, may also be useful in therapy. Upregulation of immune responses may be in the form of enhancing an existing immune response or eliciting an initial immune response. For example, enhancing an immune response through stimulating B lymphocyte antigen function may be useful in cases of viral infection. In addition, systemic vital diseases such as influenza, the common cold, and encephalitis might be alleviated by the administration of stimulatory forms of B lymphocyte antigens systemically.

Alternatively, anti-viral immune responses may be enhanced in an infected patient by removing T cells from the patient, costimulating the T cells *in vitro* with viral antigen-pulsed APCs either expressing a peptide of the present invention or together with a stimulatory form of a soluble peptide of the present invention and reintroducing the *in vitro* activated T cells into the

patient. Another method of enhancing anti-vital immune responses would be to isolate infected cells from a patient, transfect them with a nucleic acid encoding a protein of the present invention as described herein such that the cells express all or a portion of the protein on their surface, and reintroduce the transfected cells into the patient. The infected cells would now be capable of delivering a costimulatory signal to, and thereby activate, T cells in vivo.

In another application, up regulation or enhancement of antigen function (preferably B lymphocyte antigen function) may be useful in the induction of tumor immunity. Tumor cells (e.g., sarcoma, melanoma, lymphoma, leukemia, neuroblastoma, carcinoma) transfected with a nucleic acid encoding at least one peptide of the present invention can be administered to a subject to overcome tumor-specific tolerance in the subject. If desired, the tumor cell can be transfected to express a combination of peptides. For example, tumor cells obtained from a patient can be transfected ex vivo with an expression vector directing the expression of a peptide having B7-2-like activity alone, or in conjunction with a peptide having B7-1-like activity and/or B7-3-like activity. The transfected tumor cells are returned to the patient to result in expression of the peptides on the surface of the transfected cell. Alternatively, gene therapy techniques can be used to target a tumor cell for transfection in vivo.

The presence of the peptide of the present invention having the activity of a B lymphocyte antigen(s) on the surface of the tumor cell provides the necessary costimulation signal to T cells to induce a T cell mediated immune response against the transfected tumor cells. In addition, tumor cells which lack MHC class I or MHC class II molecules, or which fail to reexpress sufficient amounts of MHC class I or MHC class II molecules, can be transfected with nucleic acid encoding all or a portion of (e.g., a cytoplasmic-domain truncated portion) of an MHC class I  $\alpha$  chain protein and  $\beta_2$  microglobulin protein or an MHC class II a chain protein and an MHC class II  $\beta$  chain protein to thereby express MHC class I or MHC class II proteins on the cell surface. Expression of the appropriate class I or class II MHC in conjunction with a peptide having the activity of a B lymphocyte antigen (e.g., B7-1, B7-2, B7-3) induces a T cell mediated immune response against the transfected tumor cell. Optionally, a gene encoding an antisense construct which blocks expression of an MHC class II associated protein, such as the invariant chain, can also be cotransfected with a DNA encoding a peptide having the activity of a B lymphocyte antigen to promote presentation of tumor associated antigens and induce tumor

specific immunity. Thus, the induction of a T cell mediated immune response in a human subject may be sufficient to overcome tumor-specific tolerance in the subject.

The activity of a protein of the invention may, among other means, be measured by the following methods: Suitable assays for thymocyte or splenocyte cytotoxicity include, without limitation, those described In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan et al., eds. Greene Publishing Associates and Wiley-Interscience (Chapter 3, Chapter 7); Herrmann et al., Proc Natl Acad Sci USA 78:2488-2492, 1981; Herrmann et al., J Immunol 128:1968-1974, 1982; Handa et al., J Immunol 135:1564-1572, 1985; Takai et al., J Immunol 137:3494-3500, 1986; Takai et al., J Immunol 140:508-512, 1988; Herrmann et al., Proc Natl Acad Sci USA 78:2488-2492, 1981; Herrmann et al., J Immunol 128:1968-1974, 1982; Handa et al., J Immunol 135:1564-1572, 1985; Takai et al., J Immunol 137:3494-3500, 1986; Bowman et al., J Virology 61:1992-1998; Takai et al., J Immunol 140:508-512, 1988; Bertagnolli et al., Cell Immunol 133:327-341, 1991; Brown et al., J Immunol 153:3079-3092, 1994.

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Assays for T-cell-dependent immunoglobulin responses and isotype switching (which will identify, among others, proteins that modulate T-cell dependent antibody responses and that affect Th1/Th2 profiles) include, without limitation, those described in: Maliszewski, *J Immunol* 144:3028-3033, 1990; and Mond and Brunswick In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan *et al.*, (eds.) Vol 1 pp. 3.8.1-3.8.16, John Wiley and Sons, Toronto 1994.

Mixed lymphocyte reaction (MLR) assays (which will identify, among others, proteins that generate predominantly Th1 and CTL responses) include, without limitation, those described In: CURRENT PROTOCOLS IN IMMUNOLOGY. Coligan et al., eds. Greene Publishing Associates and Wiley-Interscience (Chapter 3, Chapter 7); Takai et al., J Immunol 137:3494-3500, 1986; Takai et al., J Immunol 140:508-512, 1988; Bertagnolli et al., J Immunol 149:3778-3783, 1992.

Dendritic cell-dependent assays (which will identify, among others, proteins expressed by dendritic cells that activate naive T-cells) include, without limitation, those described in: Guery et al., J Immunol 134:536-544, 1995; Inaba et al., J Exp Med 173:549-559, 1991; Macatonia et al., J Immunol 154:5071-5079, 1995; Porgador et al., J Exp Med 182:255-260, 1995; Nair et al., J Virol 67:4062-4069, 1993; Huang et al., Science 264:961-965, 1994; Macatonia et al., J Exp Med 169:1255-1264, 1989; Bhardwaj et al., J Clin Investig 94:797-807, 1994; and Inaba et al., J Exp Med 172:631-640, 1990.

Assays for lymphocyte survival/apoptosis (which will identify, among others, proteins that prevent apoptosis after superantigen induction and proteins that regulate lymphocyte homeostasis) include, without limitation, those described in: Darzynkiewicz et al., Cytometry 13:795-808, 1992; Gorczyca et al., Leukemia 7:659-670, 1993; Gorczyca et al., Cancer Res 53:1945-1951, 1993; Itoh et al., Cell 66:233-243, 1991; Zacharchuk, J Immunol 145:4037-4045, 1990; Zamai et al., Cytometry 14:891-897, 1993; Gorczyca et al., Internat J Oncol 1:639-648, 1992.

Assays for proteins that influence early steps of T-cell commitment and development include, without limitation, those described in: Antica et al., Blood 84:111-117, 1994; Fine et al., Cell Immunol 155: 111-122, 1994; Galy et al., Blood 85:2770-2778, 1995; Toki et al., Proc Nat Acad Sci USA 88:7548-7551, 1991.

### Hematopoiesis Regulating Activity

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A GENX protein of the present invention may be useful in regulation of hematopoiesis and, consequently, in the treatment of myeloid or lymphoid cell deficiencies. Even marginal biological activity in support of colony forming cells or of factor-dependent cell lines indicates involvement in regulating hematopoiesis, e.g. in supporting the growth and proliferation of erythroid progenitor cells alone or in combination with other cytokines, thereby indicating utility, for example, in treating various anemias or for use in conjunction with irradiation/chemotherapy to stimulate the production of erythroid precursors and/or erythroid cells; in supporting the growth and proliferation of myeloid cells such as granulocytes and monocytes/macrophages (i.e., traditional CSF activity) useful, for example, in conjunction with chemotherapy to prevent or treat consequent myelo-suppression; in supporting the growth and proliferation of megakaryocytes and consequently of platelets thereby allowing prevention or treatment of various platelet disorders such as thrombocytopenia, and generally for use in place of or complimentary to platelet transfusions; and/or in supporting the growth and proliferation of hematopoietic stem cells which are capable of maturing to any and all of the above-mentioned hematopoietic cells and therefore find therapeutic utility in various stem cell disorders (such as those usually treated with transplantation, including, without limitation, aplastic anemia and paroxysmal nocturnal hemoglobinuria), as well as in repopulating the stem cell compartment post irradiation/chemotherapy, either in-vivo or ex-vivo (i.e., in conjunction with bone marrow

transplantation or with peripheral progenitor cell transplantation (homologous or heterologous)) as normal cells or genetically manipulated for gene therapy.

The activity of a protein of the invention may, among other means, be measured by the following methods:

Suitable assays for proliferation and differentiation of various hematopoietic lines are cited above.

Assays for embryonic stem cell differentiation (which will identify, among others, proteins that influence embryonic differentiation hematopoiesis) include, without limitation, those described in: Johansson et al. Cellular Biology 15:141-151, 1995; Keller et al., Mol. Cell. Biol. 13:473-486, 1993; McClanahan et al., Blood 81:2903-2915, 1993.

Assays for stem cell survival and differentiation (which will identify, among others, proteins that regulate lympho-hematopoiesis) include, without limitation, those described in: Methylcellulose colony forming assays, Freshney, In: Culture of Hematopoietic Cells. Freshney, et al. (eds.) Vol pp. 265-268, Wiley-Liss, Inc., New York, N.Y 1994; Hirayama et al., Proc Natl Acad Sci USA 89:5907-5911, 1992; McNiece and Briddeli, In: Culture of Hematopoietic Cells. Freshney, et al. (eds.) Vol pp. 23-39, Wiley-Liss, Inc., New York, N.Y. 1994; Neben et al., Exp Hematol 22:353-359, 1994; Ploemacher, In: Culture of Hematopoietic Cells. Freshney, et al. eds. Vol pp. 1-21, Wiley-Liss, Inc., New York, N.Y. 1994; Spoonceret al., In: Culture of Hematopoietic Cells. Freshhey, et al., (eds.) Vol pp. 163-179, Wiley-Liss, Inc., New York, N.Y. 1994; Sutherland, In: Culture of Hematopoietic Cells. Freshney, et al., (eds.) Vol pp. 139-162, Wiley-Liss, Inc., New York, N.Y. 1994.

### **Tissue Growth Activity**

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A GENX protein of the present invention also may have utility in compositions used for bone, cartilage, tendon, ligament and/or nerve tissue growth or regeneration, as well as for wound healing and tissue repair and replacement, and in the treatment of burns, incisions and ulcers.

A protein of the present invention, which induces cartilage and/or bone growth in circumstances where bone is not normally formed, has application in the healing of bone fractures and cartilage damage or defects in humans and other animals. Such a preparation employing a protein of the invention may have prophylactic use in closed as well as open fracture reduction and also in the improved fixation of artificial joints. De novo bone formation

induced by an osteogenic agent contributes to the repair of congenital, trauma induced, or oncologic resection induced craniofacial defects, and also is useful in cosmetic plastic surgery.

A protein of this invention may also be used in the treatment of periodontal disease, and in other tooth repair processes. Such agents may provide an environment to attract bone-forming cells, stimulate growth of bone-forming cells or induce differentiation of progenitors of bone-forming cells. A protein of the invention may also be useful in the treatment of osteoporosis or osteoarthritis, such as through stimulation of bone and/or cartilage repair or by blocking inflammation or processes of tissue destruction (collagenase activity, osteoclast activity, etc.) mediated by inflammatory processes.

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Another category of tissue regeneration activity that may be attributable to the protein of the present invention is tendon/ligament formation. A protein of the present invention, which induces tendon/ligament-like tissue or other tissue formation in circumstances where such tissue is not normally formed, has application in the healing of tendon or ligament tears, deformities and other tendon or ligament defects in humans and other animals. Such a preparation employing a tendon/ligament-like tissue inducing protein may have prophylactic use in preventing damage to tendon or ligament tissue, as well as use in the improved fixation of tendon or ligament to bone or other tissues, and in repairing defects to tendon or ligament tissue. De novo tendon/ligament-like tissue formation induced by a composition of the present invention contributes to the repair of congenital, trauma induced, or other tendon or ligament defects of other origin, and is also useful in cosmetic plastic surgery for attachment or repair of tendons or ligaments. The compositions of the present invention may provide an environment to attract tendon- or ligament-forming cells, stimulate growth of tendon- or ligament-forming cells, induce differentiation of progenitors of tendon- or ligament-forming cells, or induce growth of tendon/ligament cells or progenitors ex vivo for return in vivo to effect tissue repair. The compositions of the invention may also be useful in the treatment of tendonitis, carpal tunnel syndrome and other tendon or ligament defects. The compositions may also include an appropriate matrix and/or sequestering agent as a career as is well known in the art.

The protein of the present invention may also be useful for proliferation of neural cells and for regeneration of nerve and brain tissue, *i.e.* for the treatment of central and peripheral nervous system diseases and neuropathies, as well as mechanical and traumatic disorders, which involve degeneration, death or trauma to neural cells or nerve tissue. More specifically, a protein

may be used in the treatment of diseases of the peripheral nervous system, such as peripheral nervous injuries, peripheral neuropathy and localized neuropathies, and central nervous system diseases, such as Alzheimer's, Parkinson's disease, Huntington's disease, amyotrophic lateral sclerosis, and Shy-Drager syndrome. Further conditions which may be treated in accordance with the present invention include mechanical and traumatic disorders, such as spinal cord disorders, head trauma and cerebrovascular diseases such as stroke. Peripheral neuropathies resulting from chemotherapy or other medical therapies may also be treatable using a protein of the invention.

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Proteins of the invention may also be useful to promote better or faster closure of non-healing wounds, including without limitation pressure ulcers, ulcers associated with vascular insufficiency, surgical and traumatic wounds, and the like.

It is expected that a protein of the present invention may also exhibit activity for generation or regeneration of other tissues, such as organs (including, for example, pancreas, liver, intestine, kidney, skin, endothelium), muscle (smooth, skeletal or cardiac) and vascular (including vascular endothelium) tissue, or for promoting the growth of cells comprising such tissues. Part of the desired effects may be by inhibition or modulation of fibrotic scarring to allow normal tissue to regenerate. A protein of the invention may also exhibit angiogenic activity.

A protein of the present invention may also be useful for gut protection or regeneration and treatment of lung or liver fibrosis, reperfusion injury in various tissues, and conditions resulting from systemic cytokine damage.

A protein of the present invention may also be useful for promoting or inhibiting differentiation of tissues described above from precursor tissues or cells; or for inhibiting the growth of tissues described above.

The activity of a protein of the invention may, among other means, be measured by the following methods:

Assays for tissue generation activity include, without limitation, those described in: International Patent Publication No. WO95/16035 (bone, cartilage, tendon); International Patent Publication No. WO95/05846 (nerve, neuronal); International Patent Publication No. WO91/07491 (skin, endothelium).

Assays for wound healing activity include, without limitation, those described in: Winter, EPIDERMAL WOUND HEALING, pp. 71-112 (Maibach and Rovee, eds.), Year Book Medical

Publishers, Inc., Chicago, as modified by Eaglstein and Menz, J. Invest. Dermatol 71:382-84 (1978).

#### Activin/Inhibin Activity

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A GENX protein of the present invention may also exhibit activin- or inhibin-related activities. Inhibins are characterized by their ability to inhibit the release of follicle stimulating hormone (FSH), while activins and are characterized by their ability to stimulate the release of follicle stimulating hormone (FSH). Thus, a protein of the present invention, alone or in heterodimers with a member of the inhibin a family, may be useful as a contraceptive based on the ability of inhibins to decrease fertility in female mammals and decrease spermatogenesis in male mammals. Administration of sufficient amounts of other inhibins can induce infertility in these mammals. Alternatively, the protein of the invention, as a homodimer or as a heterodimer with other protein subunits of the inhibin-b group, may be useful as a fertility inducing therapeutic, based upon the ability of activin molecules in stimulating FSH release from cells of the anterior pituitary. See, for example, U.S. Pat. No. 4,798,885. A protein of the invention may also be useful for advancement of the onset of fertility in sexually immature mammals, so as to increase the lifetime reproductive performance of domestic animals such as cows, sheep and pigs.

The activity of a protein of the invention may, among other means, be measured by the following methods:

Assays for activin/inhibin activity include, without limitation, those described in: Vale et al., Endocrinology 91:562-572, 1972; Ling et al., Nature 321:779-782, 1986; Vale et al., Nature 321:776-779, 1986; Mason et al., Nature 318:659-663, 1985; Forage et al., Proc Natl Acad Sci USA 83:3091-3095, 1986.

### Chemotactic/Chemokinetic Activity

A protein of the present invention may have chemotactic or chemokinetic activity (e.g., act as a chemokine) for mammalian cells, including, for example, monocytes, fibroblasts, neutrophils, T-cells, mast cells, eosinophils, epithelial and/or endothelial cells. Chemotactic and chemokinetic proteins can be used to mobilize or attract a desired cell population to a desired site of action. Chemotactic or chemokinetic proteins provide particular advantages in treatment of wounds and other trauma to tissues, as well as in treatment of localized infections. For example,

attraction of lymphocytes, monocytes or neutrophils to tumors or sites of infection may result in improved immune responses against the tumor or infecting agent.

A protein or peptide has chemotactic activity for a particular cell population if it can stimulate, directly or indirectly, the directed orientation or movement of such cell population. Preferably, the protein or peptide has the ability to directly stimulate directed movement of cells. Whether a particular protein has chemotactic activity for a population of cells can be readily determined by employing such protein or peptide in any known assay for cell chemotaxis.

The activity of a protein of the invention may, among other means, be measured by following methods:

Assays for chemotactic activity (which will identify proteins that induce or prevent chemotaxis) consist of assays that measure the ability of a protein to induce the migration of cells across a membrane as well as the ability of a protein to induce the adhesion of one cell population to another cell population. Suitable assays for movement and adhesion include, without limitation, those described in: Current Protocols in Immunology, Coligan et al., eds. (Chapter 6.12, Measurement of Alpha and Beta Chemokines 6.12.1-6.12.28); Taub et al. J Clin Invest 95:1370-1376, 1995; Lind et al. APMIS 103:140-146, 1995; Muller et al., Eur J Immunol 25: 1744-1748; Gruberet al. J Immunol 152:5860-5867, 1994; Johnston et al., J Immunol 153: 1762-1768, 1994.

#### Hemostatic and Thrombolytic Activity

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A protein of the invention may also exhibit hemostatic or thrombolytic activity. As a result, such a protein is expected to be useful in treatment of various coagulation disorders (including hereditary disorders, such as hemophilias) or to enhance coagulation and other hemostatic events in treating wounds resulting from trauma, surgery or other causes. A protein of the invention may also be useful for dissolving or inhibiting formation of thromboses and for treatment and prevention of conditions resulting therefrom (such as, for example, infarction of cardiac and central nervous system vessels (e.g., stroke).

The activity of a protein of the invention may, among other means, be measured by the following methods:

Assay for hemostatic and thrombolytic activity include, without limitation, those described in: Linet et al., J. Clin. Pharmacol. 26:131-140, 1986; Burdick et al., Thrombosis Res.

45:413-419, 1987; Humphrey et al., Fibrinolysis 5:71-79 (1991); Schaub, Prostaglandins 35:467-474, 1988.

#### Receptor/Ligand Activity

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A protein of the present invention may also demonstrate activity as receptors, receptor ligands or inhibitors or agonists of receptor/ligand interactions. Examples of such receptors and ligands include, without limitation, cytokine receptors and their ligands, receptor kinases and their ligands, receptor phosphatases and their ligands, receptors involved in cell—cell interactions and their ligands (including without limitation, cellular adhesion molecules (such as selectins, integrins and their ligands) and receptor/ligand pairs involved in antigen presentation, antigen recognition and development of cellular and humoral immune responses). Receptors and ligands are also useful for screening of potential peptide or small molecule inhibitors of the relevant receptor/ligand interaction. A protein of the present invention (including, without limitation, fragments of receptors and ligands) may themselves be useful as inhibitors of receptor/ligand interactions.

The activity of a protein of the invention may, among other means, be measured by the following methods:

Suitable assays for receptor-ligand activity include without limitation those described in: CURRENT PROTOCOLS IN IMMUNOLOGY, Ed by Coligan, et al., Greene Publishing Associates and Wiley-Interscience (Chapter 7.28, Measurement of Cellular Adhesion under static conditions 7.28.1-7.28.22), Takai et al., Proc Natl Acad Sci USA 84:6864-6868, 1987; Bierer et al., J. Exp. Med. 168:1145-1156, 1988; Rosenstein et al., J. Exp. Med. 169:149-160 1989; Stoltenborg et al., J Immunol Methods 175:59-68, 1994; Stitt et al., Cell 80:661-670, 1995.

#### **Anti-Inflammatory Activity**

Proteins of the present invention may also exhibit anti-inflammatory activity. The anti-inflammatory activity may be achieved by providing a stimulus to cells involved in the inflammatory response, by inhibiting or promoting cell—cell interactions (such as, for example, cell adhesion), by inhibiting or promoting chemotaxis of cells involved in the inflammatory process, inhibiting or promoting cell extravasation, or by stimulating or suppressing production of other factors which more directly inhibit or promote an inflammatory response. Proteins exhibiting such activities can be used to treat inflammatory conditions including chronic or acute

conditions), including without limitation inflammation associated with infection (such as septic shock, sepsis or systemic inflammatory response syndrome (SIRS)), ischemia-reperfusion injury, endotoxin lethality, arthritis, complement-mediated hyperacute rejection, nephritis, cytokine or chemokine-induced lung injury, inflammatory bowel disease, Crohn's disease or resulting from over production of cytokines such as TNF or IL-1. Proteins of the invention may also be useful to treat anaphylaxis and hypersensitivity to an antigenic substance or material.

#### **Tumor Inhibition Activity**

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In addition to the activities described above for immunological treatment or prevention of tumors, a protein of the invention may exhibit other anti-tumor activities. A protein may inhibit tumor growth directly or indirectly (such as, for example, via ADCC). A protein may exhibit its tumor inhibitory activity by acting on tumor tissue or tumor precursor tissue, by inhibiting formation of tissues necessary to support tumor growth (such as, for example, by inhibiting angiogenesis), by causing production of other factors, agents or cell types which inhibit tumor growth, or by suppressing, eliminating or inhibiting factors, agents or cell types which promote tumor growth.

#### **Other Activities**

A protein of the invention may also exhibit one or more of the following additional activities or effects: inhibiting the growth, infection or function of, or killing, infectious agents, including, without limitation, bacteria, viruses, fungi and other parasites; effecting (suppressing or enhancing) bodily characteristics, including, without limitation, height, weight, hair color, eye color, skin, fat to lean ratio or other tissue pigmentation, or organ or body part size or shape (such as, for example, breast augmentation or diminution, change in bone form or shape); effecting biorhythms or circadian cycles or rhythms; effecting the fertility of male or female subjects; effecting the metabolism, catabolism, anabolism, processing, utilization, storage or elimination of dietary fat, lipid, protein, carbohydrate, vitamins, minerals, cofactors or other nutritional factors or component(s); effecting behavioral characteristics, including, without limitation, appetite, libido, stress, cognition (including cognitive disorders), depression (including depressive disorders) and violent behaviors; providing analgesic effects or other pain reducing effects; promoting differentiation and growth of embryonic stem cells in lineages other than hematopoietic lineages; hormonal or endocrine activity; in the case of enzymes, correcting

deficiencies of the enzyme and treating deficiency-related diseases; treatment of hyperproliferative disorders (such as, for example, psoriasis); immunoglobulin-like activity (such as, for example, the ability to bind antigens or complement); and the ability to act as an antigen in a vaccine composition to raise an immune response against such protein or another material or entity which is cross-reactive with such protein.

Neural disorders in general include Parkinson's disease, Alzheimer's disease, Huntington's disease, multiple sclerosis, amyotrophic lateral sclerosis (ALS), peripheral neuropathy, tumors of the nervous system, exposure to neurotoxins, acute brain injury, peripheral nerve trauma or injury, and other neuropathies, epilepsy, and/or tremors.

#### 10 EQUIVALENTS

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From the foregoing detailed description of the specific embodiments of the invention, it should be apparent that particular novel compositions and methods involving nucleic acids, polypeptides, antibodies, detection and treatment have been described. Although these particular embodiments have been disclosed herein in detail, this has been done by way of example for purposes of illustration only, and is not intended to be limiting with respect to the scope of the appended claims that follow. In particular, it is contemplated by the inventors that various substitutions, alterations, and modifications may be made as a matter of routine for a person of ordinary skill in the art to the invention without departing from the spirit and scope of the invention as defined by the claims. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art from the foregoing description and accompanying figures. Such modifications are intended to fall within the scope of the appended claims.

PCT/US00/08621

WO 00/58473

Table 1

RF#	IRF# Internal Identification	Protein similarity	Protein domain	Protein (Classification	Calls or Tissues in which Gene is Expressed
i i	13076366 (1, 2)	Novel Protein sim. GBank gil4691395 emb CAB41562.11- (AL049727) putative large secreted protein [Streptomyces coelicolor]			264636
	80248091 (3, 4)	Novel Protein sim. GBank gi/2829506/splP71559/SUCC_MYCTU - SUCCINYL-COA SYNTHETASE BETA CHAIN (SCS-BETA)	Contains protein domain (PF00549) - UNCLASSIFIED CoA-ilgases		264907, 264600, 264602, 264762, 264769, 264689, 264638, 264567
	80415924 (5, 6)				264910, 264604, 264634, 264905, 264635, 264691, 264907, 264692, 264629
	82018837 (7, 8)				264908, 264909, 264760, 264628, 264635
	79970035 (9. 10)				2279002, 264563
	79842462 (11, 12)		Contains protein domain (PF00127) - UNCLASSIFIED Copper binding proteins, plastocvanin/azurin family		264908
·	85515576 (13, 14)	Novel Protein sim. GBank gil4415926 gb AAD20157  - (AC006282) unknown protein [Arabidopsis thallana]		UNCLASSIFIED	20281099, 35696052, 264508, 264509, 264905, 264906, 264906, 264908, 264909, 264511, 265009, 264512, 265009, 264910, 264595, 264596, 264761, 265009, 264761, 264762, 264693, 264762, 264693, 26489, 35695917, 264690, 264692, 264693, 33657109, 264628, 264630, 264631, 264632, 264634, 264636, 264631, 264631, 264634, 264636, 264564, 264566, 264666,
_	56924278 (15, 16)	Novel Protein slm. GBank gi 585562 sp 006456 NIRB_KLEPN - NITRITE pern ictae Anarowy i ABGE SHRINIT		reductase	264907
	79394457 (17, 18)			UNCLASSIFIED	265007, 265019, 263972
₽	79556459 (19, 20)			UNCLASSIFIED	264906
	20414027 (21, 22)				264605
12	94141210 (23, 24)	Novei Protein sim. GBank gij3878145 emb CAA99871  - (Z75543) similar to potassium channel protein [Caenomabditis elegans]			264259, 265007, 83373044
5	20750551 (25, 26)	1		UNCLASSIFIED	264556, 264557, 264564
4	95105114 (27, 28)	Novel Protein sim. GBank gi[2832781]emb CAA12645  - Contains pr (AJ225805) inward potassium channel alpha subunit [Egeria Ank repeat densa]	Contains protein domain (PF00023) - potassium_channei a Ank repeat		35696288, 35696052, 264510, 35695917, 264691, 264628, 35696423, 264555, 264558, 264559, 83373044
15	20458307 (29, 30)	Novel Protein sim. GBank gij1710791 sp Q10234 RT05_SCHPO - PROBABLE MITOCHONDRIAL 40S RIBOSOMAL PROTEIN S5	Contains protein domain (PF00333) - inbosomalprot Ribosomal protein S5		264604
9	20760356 (31, 32)				264555

Ŀ	1000000				
<u> </u>	20252144 (33, 34)		Contains protein domain (PF00449) Urease		264600
18	80246804 (35, 36)	Novel Protein sim. GBank gi 2281102 (AC002333) - SF16 isolog (Arabidopsis thatiana)			29331827, 264555, 264557, 264638, 264558
19	80076624 (37, 38)			UNCLASSIFIED	22278996, 264907, 264910, 264600, 264693
20	20724558 (39, 40)			transport	264602
		gi[2506112]sp[P43672]UUP_ECOLI - ABC TRANSPORTER ATP-BINDING PROTEIN UUP			
21	80417554 (41, 42)	Novel Protein sim. GBank		UNCLASSIFIED	22278995, 264906, 265008, 265010, 265011,
		gil1730203 sp P50442 GATM_RAT - GLYCINE	.:		264602, 264605, 264766, 264688, 21906764,
		AMIDINOTRANSFERASE PRECURSOR (L.			264691, 18108376, 264636, 18108387,
		ARGININE:GLYCINE AMIDINOTRANSFERASE) [TRANSAMIDINASE] (ATI			264486
22	11705858 (43, 44)				264685
23	80419176 (45, 46)	Novel Protein sim. GBank gij1877329jembįCAB070771 -	Contains protein domain (PF00441) - dehydrogenase	dehydrogenase	264488, 264907, 264909, 264600, 264602,
		(292771) fadE25 [Mycobacterium tuberculosis]	Acyl-CoA dehydrogenase		264603, 264605, 264682, 264766, 32833986, 364836, 364486
24	20291697 (47, 48)				264600
25	80253774 (49, 50)				264593
1	80255394 (51, 52)			UNCLASSIFIED	22278996 56182435 265018 264566
27	80235795 (53, 54)	Novel Protein sim. GBank gil4808369lemblCAB42783.11 -	Contains protein domain (PE00253) - Inhosomalinet	rihosomatorot	18108370 35696423 264635 264555
			Ribosomal protein S14p/S29e		
		[Streptomyces coelicolor]			
82	79483561 (55, 56)			UNCLASSIFIED	264638
59	82448765 (57, 58)	Novel Protein sim. GBank	Contains protein domain (PF00365) - kinase	kinase	264601, 284762, 264766, 264769, 264636
	<u>.</u>	gij3122290jspj008333jK6PF_STRCO - 6- PHOSPHOFRUCTOKINASE (PHOSPHOFRUCTOKINASE) (PHOSPHOHEXOKINASE) (ATP-PFK)	Phosphofuctokinase		
90	79199333 (59, 60)			UNCLASSIFIED	264908, 265019, 264687, 21906764,
31	19848158 (61, 62)			UNCLASSIFIED	21900100
32	82449495 (63, 64)	Novel Protein sim. GBank gij3560504 (AF027770) -		UNCLASSIFIED	264905, 264605, 264762, 264766, 264687,
Ę	100 000 000	unknown (Mycobacterium smegmatis)			264689
3	79582628 (65, 66)	Novel Protein sim. GBank gi 2129003 pir  G64507 - hypothetical protein MJ1665 - Methanococcus jannaschii		UNCLASSIFIED	264687
<u>స</u> ్త	87467657 (67, 68)			UNCLASSIFIED	60432289, 264600, 264602, 264760. 18108357, 264769, 285020, 264691
35	95005170 (69, 70)	Novel Protein sim. GBank gij5420387[emb CAB46679.1] - (AJ243459) proteophosphoglycan [Leishmania major]		UNCLASSIFIED	264600, 264687, 264558, 264639
ဗ္က	19642042 (71, 72)	Novel Protein sim. GBank gi]3287739[sp P73538]BIOB_SYNY3 - BIOTIN SYNTHASE (BIOTIN SYNTHETASE)		synthase	264566
37	20369215 (73, 74)	Novel Protein sim. GBank gi[2313134]gbJAAD07126.11 - (AE00527) delta-1-pyrroline-5-carboxylate dehydrogenase [[Helicobacier pylori 26695]		dehydrogenase	264603

264605	264905, 264906, 264907, 66712502, 264908, 264909, 264909, 264511, 265009, 264910, 55812038, 264758, 265011, 264762, 264682, 264763, 264764, 264766, 265022, 264693, 264628, 264631, 264634, 264555, 264582, 264581, 264581, 264582, 264582, 264582, 264582, 264882	264592	264591, 35695917	264602	264605	264769, 264638	264769, 264510, 264508	264566	264689	18108385, 264635, 264828	264603	264508, 264603, 264769, 264689, 264635, 264558, 284486	264593, 18108387	264634	264762	264630, 264909, 264766	29331824, 264102, 265018, 18108376	264604	264557
reductase	complement	UNCLASSIFIED		UNCLASSIFIED	synthase	synthase	UNCLASSIFIED	ngf	UNCLASSIFIED		UNCLASSIFIED		UNCLASSIFIED	struct	UNCLASSIFIED	UNCLASSIFIED			- UNCLASSIFIED
	Contains protein domain (PF00207) - complement Alpha-2-macroglobulin family					Contains protein domain (PF00958) - synthase GMP synthase C terminal domain													Contains protein domain (PF00076) - UNCLASSIFIED RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)
Novel Protein sim. GBank gij3805970jembjCAA06231 - (AJ004933) periplasmic nitrate reductase, large subunit IRhadonsendomonas so i	Novel Protein sim. GBank gij1929449 (L63543) - endodermin [Xanopus laevis]		Novel Protein sim. GBank gil854065jembjCAA58337j -		Novel Protein sim. GBank gij3820584 (AF086791) - carbamoyiphosphate synthelase large subunit (Zymomonas mohilie)	Novel Protein sim. GBank gi2494784 sp Q50729 GUAA_MYCTU - GMP SYNTHASE (GLUTAMINE-HYDROLYZING) (GLUTAMINE AMIDOTRANSFERASE) (GMP SYNTHETASE)							Novel Protein sim. GBank gi[3411177 (AF076240) - MocC Rhizoblum teguminosarum bv. viciae]	Novel Protein sim. GBank gij3914992 sp QZ6264 SM41_HEMPU - 41 KD SPICULE MATRIX PROTEIN PRECLIRSOR (HSM41) (HPSMC)	Novel Protein sim. GBank gij3980411 (AC004561) - putative profine-rich protein [Arabidopsis thaliana]		Novel Protein sim. GBank gij 1633572 (U52064) - Herpesvirus salmiri ORF73 homolog [Kaposi's sarcoma- associated hemse-like virus]	Novel Protein sim. GBank gil4321580 gb AAD15785  - (AF050114) alqinate byase [Pseudomonas sp. W7]	
20466334 (75, 76) N (A	94300715 (77, 78) el	20635625 (79 80)		T	20467069 (85, 86)	13085297 (87, 88) N	39384711 (89, 90)	95003398 (91 92)	11698624 (93 94)	79407218 (95 96)	21659844 (97, 98)	80503996 (99, 100)	80255569 (101, 102)	79208528 (103, 104)	36996970 (105, 106)	79570897 (107, 108)		8758408 (111, 112)	11223386 (113, 114)
38	86	Ç	T	Τ	£	4.	45	å,	2 5		64	S <sub>S</sub>	51	25	S	Z,	55	28	25

9	104002500		11 10000000		C300000 01000100 020100 12200101
3	91427309 (113, 119)	100ver Frudelin Sim. CBdain gil5616074[gb]AAD45616.1[AF06194 - (AF061943) protate-		Dep. III	66712502, 264909, 265008, 265010, 265011.
		derived STE20-like kinase PSK [Homo saplens]			264681, 29148784, 35695917, 60170615,
					264691, 264692, 264693, 18108374,
					35696423, 56182323, 60432113
59	80077371 (117, 118)	Novel Protein sim. GBank	Contains protein domain (PF00953) - Irransferase	transferase	264600, 264689, 264638
		gij1172920jspjP45830jRFE_MYCLE - PUTATIVE	Glycosyl transferase		
		UNDECAPRENYL-PHOSPHATE ALPHA-N-			
8	12958341 (119, 120)				264689
91	80426808 (121, 122)	Novel Protein sim. GBank gil1710216 (U79260) - unknown		glycoprotein	264768
		[Homo sapiens]			
62	13504966 (123, 124)				264630
63	16474553 (125, 126)			UNCLASSIFIED	265019
2	20724578 (127, 128)	Novel Protein sim. GBank gij420945[pirj]A47041 -		UNCLASSIFIED	264602
		fransposase homolog (insertion element ISAE1) -			
		Alcaligenes eutrophus			
65	79326308 (129, 130)		Contains protein domain (PF00224) - kinase	kinase	264563
		gij3122312jsp OD6134 KPYK_MYCTU - PYRUVATE   KINASE (PK)	Pyruvate kinase		
99	46854384 (131 132)	Novel Protein cim CRank aligo28723lembiCA6222101.		transmit	2227R006 26455R
}	(201 (101) 1021 2021	(AL034355) putative ABC transporter (Strentomyces			
		coeficolor			
29	78952543 (133, 134)	Novel Protein sim. GBank		dehydrogenase	265021
		gi[231985 sp P30234 DHA MYCTU - ALANINE			
		DEHYDROGENASE (40 KD ANTIGEN)			
88	79817382 (135, 136)				264909
69	79841764 (137, 138)			UNCLASSIFIED	264908
2	79871329 (139, 140)				264906, 264908
71	65897458 (141, 142)			UNCLASSIFIED	264602, 265021
72	87734977 (143, 144)	Novel Protein sim. GBank gil4415926lgblAAD201571 -		UNCLASSIFIED	264488, 264905, 264906, 264907, 264908.
		(AC006282) unknown protein [Arabidopsis thaliana]			264511, 265008, 264910, 264758, 87168474,
					264682, 264766, 264686, 264689, 35695917,
					265021, 60170615, 264691, 33657023.
					264692, 264693, 264629, 264631, 264639,
					22279000
2	80025241 (145, 146)			UNCLASSIFIED	60424179, 264508, 264908, 265007, 264603.
ŀ	.000				264667, 264669, 264692, 18108387
٠	2037/410 (147, 148)			UNCLASSIFIED	264605
75	11819032 (149, 150)	Novel Protein sim. GBank gij2853098jembjCAA16914j -		UNCLASSIFIED	264689
		(ALUZ 1 / 0 / ) Vacuotar protein soning (Scrizosaccharomyces (pombe)	vo.		
92	95105303 (151, 152)	Novel Protein sim. GBank gij4468811 jembjCAB38212 j		UNCLASSIFIED	83373044, 264906, 264557
		(AL035601) putative protein [Arabidopsis thaliana]			
<u>,                                    </u>	10144/18 (153, 154)	Novel Protein sim. GBank gij854065jemb[CAA58337] - (X83413) U88 [Human herpesvirus 6]		UNCLASSIFIED	264563
78	8758258 (155, 156)			UNCLASSIFIED	264604

62	94140190 (157, 158)	Novel Protein sim. GBank gil5689453 dbj BAA83010.1  -	Contains protein domain (PF00169) -		35696286, 22278998, 29331822, 29331824,
					. 29331823, 29331827, 264903, 264906, 264906, 264907, 66712502, 264908, 264909, 265008, 265009, 265009, 265009, 265009, 2649009, 2649000, 26490000, 2649000, 2649000, 2649000, 2649000
			•		33109954, 265017, 265018, 264288, 264768.
					56181562, 21906765, 21906769, 29148784,
					265020, 264690, 264691, 264692, 264693.
					60431528, 35696423, 264631, 264632,
					264634, 264636, 264639, 83373044, 264564, 1264566, 284587
80	82314840 (159, 160)			UNCLASSIFIED	264769, 264601, 265006, 264910, 264604.
					264605, 264634, 264635, 264905, 264762,
					264637, 264592, 264628, 264907, 264691,
2	20467247 (161, 162)	Novel Protein sim. GBank		reductase	264605
	•	dil1723442lspi010258lYD2A SCHPO - HYPOTHETICAL			
		69.0 KD PROTEIN CS6F8.10 IN CHROMOSOME I			
82	16331388 (163, 164)	Novel Protein sim. GBank gi 2895866 (AF045770) -		dehydrogenase	264567
		methylmatonate semi-aldehyde dehydrogenase [Oryza			
		sativa]			
£	(94741180 (165, 166)	Novel Protein sim. GBank gij3402673 (AC004697) -		UNCLASSIFIED	264488, 264508, 264509, 264905, 264908.
		unknown protein (Arabidopsis thaliana)			264909, 264511, 264591, 264593, 264594,
					264595, 264596, 264758, 264603, 264760,
					264681, 18108351, 264762, 264682, 264764.
					264684, 264766, 264686, 264632, 264637,
8	80355375 (167, 168)	Novel Protein sim. GBank		francont	264337, 264636, 264639, 18108383, 264366
		gij1173364 spIP45380 SAT1_RAT - SULFATE ANION			264910, 284760, 264763, 264764, 264768
		TRANSPORTER 1 (CANALICULAR SULFATE			264768, 264769, 35695855, 264636, 264637
		TRANSPORTER) (SULFATE/CARBONATE ANTIPORTER)			
92	80499600 (169, 170)	Novel Protein sim. GBank gij2120998ipirijS70682 -		transferase	264605, 264762, 264687, 264769, 18108374,
		glycosyltransferase homolog - Bordetella pertussis			264636, 264486
8	39559043 (171, 172)	Novel Protein sim. GBank gij3256023jembjCAA17228.1j -			264910
		(ALUZ1897) hypothetical protein Rv1112 [Mycobacterium			
2	13856808 (173, 174)				
	1202000 (113, 114)			UNCLASSIFIED	1264093

95344718 (175, 176) 80077389 (177, 178) 82115999 (179, 180) 78906950 (181, 182) 79554871 (183, 184) 80496778 (185, 186)

1	94322125 (191, 192)	Novel Protein sim. GBank gil4589560 dbj BAA76802.1  -		UNCLASSIFIED	22278995, 22278999, 264259, 29331822,
					264906, 264907, 264908, 264909, 265007, 265008, 284910, 265009, 264593, 265010, 265017, 264604, 265019, 18108351, 264288, 264766, 264768, 264769, 21906765, 24906762, 2565020, 264692, 33657182, 35695763, 264628, 264629,
- 6					18108379, 264631, 264638, 18108381, 264559, 18108382, 83373044, 22279002
	79605200 (193, 194)	Novel Protein sim. GBank gil4583559 emb CAB40388.1  - (AJ005255) OxyR (Erwinla chrysanthem)			264508
	79427000 (195, 196)	Novel Protein sim. GBank gi 1001693 db  BAA10430  - (D64002) hypothetical protein (Synechocystis sp.)			264909
	20466524 (197, 198)	Novel Protein sim. GBank gij1169479jspjP43925jEFG_HAEIN - ELONGATION FACTOR G (EF-G)			264605
	79640113 (199, 200)				264693
	80203298 (201, 202)	Novel Protein sim. GBank gil480897 pir  S37485 - gene msg1 protein - mouse		UNCLASSIFIED	265020, 264102, 263972
	20467259 (203, 204)	Novel Protein sim. GBank gi 2894166 emb CAA11773.1 - (AJ223998) PCZA361.18 [Amycolatopsis orientalis]		synthase	264605
	20466368 (205, 206)	H.	Contains protein domain (PF00271) - Inelicase Helicases conserved C-terminal domain	helicase	264605
	80247572 (207, 208)	Novel Protein sim. GBank gil854065 emb CAA58337  - (X83413) U88 [Human herpesvirus 6]		UNCLASSIFIED	264591, 264595, 264602
i	79605206 (209, 210)	Novel Protein sim. GBank gi 1685117 (U70770) - furrowed [Drosophila metanogaster]	Contains protein domain (PF00084) - complement Sushi domain (SCR repeat)		264508
	28382058 (211, 212)	Noveł Protein sim. GBank  gi[1705505]sp[P54729]BS4_MOUSE - BS4 PROTEIN	F00627) -	UNCLASSIFIED	264511, 265009
	80057791 (213, 214)	Novel Protein sim. GBank gij4887229[gbjAAD32244.1 AF15075 - (AF150755) microtubule-actin crosslinking factor [Mus musculus]		ATPase_associated	ATPase_associated   29331624, 264591, 21906754, 265019
	80237936 (215, 216)	Novel Protein sim. GBank gi 2635771 emb CAB15264  - (299120) similar to ABC transporter (ATP-binding protein) {Bacillus subtilis}	Contains protein domain (PF00005) - transport ABC transporter	transport	18108374, 35695917, 22278996, 264113, 264600, 264602, 264603, 285017, 264910. 264906, 264636, 264766
1	95194148 (217, 218)	Novel Protein sim. GBank gil2330791 emb CAB11265  - (298601) carboxypeptidase s precursor  Schizosaccharomyces pombe]		UNCLASSIFIED	264758, 264603, 264630, 264636, 264637
. 1	79582823 (219, 220)				264687
	39565458 (221, 222)				264564
1	78856038 (223, 224)				264908
1	80502101 (227, 228)			UNCLASSIFIED	265007 264769
1	100306101 (661, 660)			UNCLASSIFIED	704/09

52645156, 52645080, 33656970, 264592, 21908754, 27486264, 18108379, 35696423, 264635, 52644332, 18108382	284905, 284906, 284908, 284909, 284695, 284909, 264910, 284768, 285010, 284763, 284682, 284768, 284768, 284768, 284768, 284768, 284768, 284693, 3365709, 284638, 284631, 284632, 284638, 284639, 56526486, 284565, 284566	264639, 264693	263974	22278996, 29331824, 60432289, 263007, 60433438, 264605, 18108351, 264769, 264689, 265020, 284534, 27486261, 264558, 83373044, 18108385, 264564	264603	264595	264605	264604	18108394, 18108397, 265006, 265007, 265008, 265010, 265011, 18108355, 18108379, 18108380, 18108384	264508, 264906, 265009, 264596, 22278002	264511	264605	264907, 264909, 264910, 264600, 264601, 264603, 264605, 18108351, 264693, 264557	35696423, 35695855, 264600, 264602, 264603, 264604, 264605, 264508, 264506, 264564, 264628, 264682, 264565, 264683	264634	18108376, 264769, 29331826, 264689, 22278996, 265021, 284600, 264511, 284601, 264602, 264605, 264905, 264836
UNCLASSIFIED			UNCLASSIFIED	UNCLASSIFIED	kinase		transferase		synthase	misc_channel		UNCLASSIFIED	phosphalase	UNCLASSIFIED		- transport It
			Contains protein domain (PF00170) - UNCLASSIFIED bZIP transcription factor				Contains protein domain (PF00398) - Iransferase Ribosomal RNA adenine dimethylases			Contains protein domain (PF00060) - misc_channel Ligand-gated ion channel			Contains protein domain (PF00072) - phosphatase Response regulator receiver domain	Contains protein domain (PF00270) - UNCLASSIFIED DEAD/DEAH box helicase		Contains protein domain (PF00528) - transport Binding-protein-dependent transport systems inner membrane component
Novel Protein sim. GBank gi[2246532 (U93872) - ORF 73, contains large compare repeat CR 73 [Kaposi's sarcoma-				Novel Protein sim. GBank gij732526 (U22327) - alpha2(IV) collagen [Caenorhabditis elegans]	Novel Protein sim. GBank gi[2131219 pir  S50157 - cyclindependent kinase chain SRB10 - yeast (Saccharomyces		Novel Protein sim. GBank gi 2052147 emb CAB08137  - (294752) ksgA [Mycobacterium tuberculosis]		Novel Protein sim. GBank git283385[sp[043134]UGST_SORBI - GRANULE-BOUND gi YCOGEN (STARCH) SYNTHASE PRECURSOR	Novel Protein sim. GBank gild75542 (U08255) - glutamate recentre delta-1 subunit (Rattus novecicus)	Novel Protein sim. GBank gi 5102785 emb CAB45200.1  - (A.079308) putative transcriptional regulator [Streptomyces coelicidati		Novel Protein sim. GBank gi[130120]sp[P23620]PHOB_PSEAE - PHOSPHATE gi[130120] TRANSCRIPTIONAL REGULATORY PROTEIN	Protein sim. GBank 493[sp[P38036]YG CD PROTEIN IN IAI	Novel Protein sim. GBank gij854065jembjCAA58337j -	Novel Protein sim. GBank gij 1076038 piri S54860 - ABC transporter PstC-2 chain - Mycobacterlum tuberculosis
80251003 (229, 230) N	81288689 (231, 232)	78636695 (233, 234)	80222170 (235, 236)	91013071 (237, 238)	8756491 (239, 240)	24.2)	20457620 (243, 244)	1076 370 (376)	79104017 (247, 248)	87797986 (249, 250)	56701283 (251, 252)	196 6367 7965 364)	80248473 (255, 256)	95290543 (257, 258)	80085583 (259, 260)	94995022 (261, 262)
215	91.16	117 7	T	119	120	Т	122	Т	124	125	126	Τ	128	128	130	131

1	110887692 (263 264)				000100
133	94630883 (265, 268)	Novel Protein eim GBank gil 1877340tembl CABOZOEBI	Contains acatain domain (BEO0380)	Conference	204030
1			Carbamoyl-phosphate synthase (CPSase)	) colonial c	104200, 104000
<u>¥</u>	79834660 (267, 268)	Novel Protein sim. GBank gil4585838jembiCAB40932.1j - (AL049630) putative NADH dehydrogenase [Streptomyces coellooter]		dehydrogenase	264905, 264605, 265021
135	19885057 (269, 270)	Novel Protein sim. GBank gi]1460074[emb]CAB01049] - (277250) hypothetical protein Rv2568 [Mycobacterium fuberculosis]			264634
136	79846083 (271, 272)	Novel Protein sim. GBank gij2125896jembjCAA73511  - (Y13070) folypolyglutamate synthase [Streptomyces coellcolor]		synthase	264508
137	79619770 (273, 274)				264683, 264685, 264686, 264691, 264692, 264693
138	79635971 (275, 276)			UNCLASSIFIED	18108374, 18108385, 33657109, 33657182, 265010, 22278998, 265008, 265007, 265008, 265009, 264693
139	86688076 (277, 278)	Novel Protein sim. GBank gi 5689912 emb CAB52075.1  - (AL109732) putative mutase [Streptomyces coelicolor A3(2)]	Contains protein domain (PF01817) - dehydrogenase Chorlsmate mutase	dehydrogenase	22278996, 265007, 264910, 60433356, 265010, 264602, 264605, 264768, 264688, 264769, 264693, 32833986, 18108374, 18108387
<del>5</del>	79825759 (279, 280)			UNCLASSIFIED	284908
Ξ	20700094 (281, 282)				264600
142	80028104 (283, 284)	Novel Protein sim. GBank gij3581916 emb CAA20855  - (AL031545) mutS family DNA mismatch repair protein [Schizosaccharomyces pombe]		nuclease	284602, 265017
143	11072274 (285, 286)			UNCLASSIFIED	264600
<del>4</del>	95009102 (287, 288)	Novel Protein sim. GBank gi 3334127 sp P97303 BAC2_MOUSE - TRANSCRIPTION REGULATOR PROTEIN BACH2 (BTB AND CNC HOMOLOG 2)			263978, 264600, 264910, 264632, 264508, 264563, 264564, 264591, 264556, 264908, 264629, 264639
145	80027058 (289, 290)	Novel Protein sim. GBank gij3757569 emb CAA21315  - (AL031863) 1-evidence=predicted by content; 1-melhod=genefinder;084; 1-melhod_score=66,31; 1-evidence_end [Drosophila melanogaster]		UNCLASSIFIED	22278996, 264602
146	13085662 (281, 292)	Novel Protein sim. GBank gif140807 sp P24536 Y121_BURCE - INSERTION ELEMENT IS402 HYPOTHETICAL 24 KD PROTEIN	Contains protein domain (PF01675) - Transposase		264887
147	94320366 (293, 294)	Novel Protein sim. GBank gij2827608 emb CAA16663  - (AL021646) uvrD2 [Mycobacterium tuberculosis]		helicase	264905, 264906, 264909, 264510, 265009, 60433356, 264600, 284601, 264604, 264605, 264687, 264769, 18108365, 65274791, 18108387
84	80248804 (295, 296)	Novel Protein sim. GBank gi[2916947 emb CA417585  - (AL021999) hypothetical protein Rv0986 [Mycobacterium [tuberculosis]		transport	265009, 265010, 264600, 264602, 264603, 264604, 264605, 264693, 33657109, 264636

149	80249373 (297, 298)	Novel Protein sim. GBank gil1723073lsplQ11040 Y081 MYCTU - HYPOTHETICAL	Contains protein domain (PF00005) - Iransport ABC transporter	Iransport	265010, 264600, 284601, 264603, 264604, 27486265, 264636
150	20294748 (299, 300)	Novel Protein sim. GBank gij3724125 emb CAA11905  - (AJ224340) maltosephosphorylase [Lactobacillus			264600
		sanfrancisco]			
151	20726398 (301, 302)	Novel Protein sim. GBank	Contains protein domain (PF01676) - UNCLASSIFIED Metalinenzyme superfamily	UNCLASSIFIED	264602
		PHOSPHOPENTOMUTASE			
		(PHOSPHODEOXYRIBOMUTASE)			
152	95002877 (303, 304)			peptidase	264602
		g  Z49/95Z sp  P5555Z 141M_KHISN - HYPOTHETICAL  HYDROLASE/PEPTIDASE Y4TM			
153	80256665 (305, 306)	Novel Protein stm. GBank		UNCLASSIFIED	264593
		gij3123021 splQ90508 VIT1_FUNHE - VITELLOGENIN I			
		PRECURSOR (VTG I) (CONTAINS: LIPOVITELLIN 1 (LV1); PHOSVITIN (PV): I IPOVITE! (IN 2 d V2))			
25	82305966 (307, 308)	(1)-			264910, 264762, 264691, 264634
155	20429859 (309, 310)	Novel Protein sim GBank gil419697Iniril.JN0443 -	Contains protein domain (PE00140) -	manolymerase	264605
<u> </u>		transcription initiation factor sigma homolog hrdB -	Sigma-70 factor		
		Streptomyces aureofaciens	1		
156	39564742 (311, 312)	Novel Protein sim. GBank gi 628710 pir  S41739 -		UNCLASSIFIED	264565
		hypothetical protein - Escherichia coli			
157	10358887 (313, 314)	Novel Protein sim. GBank gij3695013 (AF052586) - CtrA	Contains protein domain (PF00142) - hydrolase	hydrolase	264691
		(Pseudomonas aeruginosa)	4Fe-4S iron sulfur cluster binding		
			proteins, NifH/frxC family		
2	79761936 (315, 316)	Novel Protein sim. GBank gil 1073072 pir  C55543 · cmaU  protein - Pseudomonas syringae pv. syringae		UNCLASSIFIED	264905
159	78890376 (317, 318)			UNCLASSIFIED	265008
160	11075119 (319, 320)		Contains protein domain (PF00400) -		264605
			WD domain, G-beta repeat		
161	80055007 (321, 322)	Novel Protein sim. GBank	Contains protein domain (PF00327) - ribosomalprot	ribosomaíprot	22278996, 264600, 264603, 35695917,
			Ribosomal protein L30p/L7e		32833986, 35696423, 264636
162	80016371 (323, 324)	Novel Protein sim. GBank gij5304869jembjCAB46028.11 -	Contains protein domain (PF00097) - interleukin	interleukin	264112, 264532, 22279002
		ליביסים מייסים (היסים היסים ליסים ליסים היסים ליסים היסים ליסים היסים ליסים היסים ליסים היסים ליסים היסים ליסים	(inger)		
163	11692306 (325, 326)			UNCLASSIFIED	264639
<u>1</u>	80077902 (327, 328)			UNCLASSIFIED	264905, 264907, 264600
忘	10856067 (329, 330)				264691
<u>8</u>	88095003 (331, 332)	Novel Protein sim. GBank gij2661691 jembjCAA15795 j - (AL 009204) putative protease (Streptomyres coefficilor)		UNCLASSIFIED	264605, 264486
167	16395460 (333, 334)	Novel Protein sim. GBank gil4416478 qb AAD20378  -		UNCLASSIFIED	265010
		(AF125999) transposase [Mycobacterium avium]			
20	80079362 (335, 336)	Novel Protein sim. GBank gi 76177 pir  QQECFT - hypothetical 38.8K protein (fisl 5' region) - Escherichia coli			264600
169	80239581 (337, 338)				264556, 264557, 264558, 264559

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	79612364 (339, 340)				2007.00
Ē	95293073 (341, 342)	Novel Protein sim. GBank	Contains profein domain (PE01810)		264505 264604
		gij 40888 spp27847 YIGK_ECOLI - HYPOTHETICAL 15.4 KD PROTEIN IN RECQ-PLDB INTERGENIC REGION (F138)			*CO+OA), KO+OCO+
172	37797007 (343, 344)		Contains protein domain (PF00528) - transport Binding-protein-dependent transport systems inner membrane component	transport	264769
173	57529660 (345, 346)	Novel Protein sim. GBank gil 132854 spjP02387 RL2_ECOLI Contains protein domain (PF00181) - ribosomalprot - 50S RIBOSOMAL PROTEIN L2 Ribosomal Proteins L2	Contains protein domain (PF00181) - Ribosomal Proteins L2	ribosomalprot	264769
174	95293078 (347, 348)	Novel Protein sim. GBank gi 1881350 dbj BAA19377 - (AB001488) PROBABLE TRANSPORT PROTEIN, SIMILAR TO ANTIBIOTIC TRANSPORT-ASSOCIATED PROTEIN ACTII IN STREPTOMYCES COELICOLOR, [Bacillus subtilis]		transport	264510, 264593, 264602, 264603, 264605, 264762, 284693
175	79756270 (349, 350)	Novel Protein sim. GBank gil2072722[emb]CAB08326] - (295121) manA [Mycobacterium tuberculosis]		isomerase	264565
176	80066896 (351, 352)	Novel Protein sim. GBank gil1055198 (U40187) - similar to PIR:A41724 chicken LD (limb deformity) gene product and to formin; also P-rich region similar to collagen [Caenorhabditis elegans]		UNCLASSIFIED	264907, 264910, 264681, 264558
177	86684852 (353, 354)	Novel Protein sim. GBank gil2326738 emb CAB10952  - (298288) hypothetical protein Rv1695 [Mycobacterfum tuberculosis]	Contains protein domain (PF01513) - UNCLASSIFIED Domain of unknown function	UNCLASSIFIED	264768, 60424179, 264687, 264688, 264769, 29331828, 60432289, 18108376, 264689, 18108387, 32833986, 22278998, 265020, 264600, 264601, 264602, 264603, 264604, 264605, 264638, 264586, 264564, 264564, 264638, 264486, 60433356, 264866
178 2	79559526 (355, 356)	Novel Protein sim. GBank gil 1906596 (U81788) - kinesin-73 [Drosophila melanogaster]		struct	264693, 33657109, 264635
179	20263112 (357, 358)			UNCLASSIFIED	264563
180	80488958 (359, 360)	Novel Protein sim. GBank gij1169387jspjP45256jDNAB_HAEIN - REPLICATIVE DNA HELICASE		helicase	264769
181	79585369 (361, 362)	Novel Protein sim. GBank gij3170615 (AF059485) - DOC4 [Mus musculus]		UNCLASSIFIED	21906767, 264635, 264639, 18108384
182	80577899 (363, 364)			UNCLASSIFIED	264259, 35696052, 56182435, 264511, 265018, 33657109, 264555, 264566
183	11614017 (365, 366)	Novel Protein sim. GBank gi 1076627 pir  S54172 - inorganic pyrophosphatase (EC 3.6.1.1) - common tobacco		UNCLASSIFIED	264690
<u>\$</u>	10174167 (367, 368)	Novel Protein sim. GBank gil4371280 gb AAD18138  - (AC006260) hypothetical protein (Arabidopsis thaliana)		UNCLASSIFIED	264510

185	21660822 (369, 370)			UNCLASSIFIED	264604
		(AL022304) putative mma transport regulator [Schizosaccharomyces pombe]	٠.		
186	80070329 (371, 372)	Novel Protein sim. GBank gi 2829802 sp P94408 YCLF_BACSU - HYPOTHETICAL		transport	264595
187	B0186611 (373 374)	53.3 KD PROTEIN IN SFP-GERKA INTERGENIC REGION			
	20160011 (373, 374)			UNCLASSIFIED	264369
8	20464942 (375, 376)	Novel Protein sim. GBank gij3150260jembjCAA19179j - [AL023634) cyclin [Schizosaccharomyces pombe]		kinase	264605
189	82338215 (377, 378)	Novel Protein sim. GBank gi[2145853 pirj \$72938 - hftx		UNCLASSIFIED	35696052, 264602, 264605, 264762, 264689,
		process - mycobacterium leprae			35695917, 18108370, 18108372, 264638, 264585
190	80088821 (379, 380)		Contains protein domain (PF00205) - synthase	synthase	284563
		(AB001488) SIMILAR TO PYRUVATE OXIDASE AND ACETOLACTATE SYNTHASE. (Bacillus subtilis)	Thiamine pyrophosphate enzymes		
181	88095012 (381, 382)	Novel Protein sim. GBank	Contains protein domain (PF00254) - isomerase	isomerase	264508 264604 264605 264769 264555
		gil120226 spiP28725 FKBP_STRCH - FK508-BINDING			
		PROTEIN (PEPTIDYL-PROLYL CIS-TRANS ISOMERASE)   (PPIASE) (ROTAMASE)			
192	16333379 (383, 384)				251557
193	79910127 (385, 386)				SEASON SEASON
194	20464949 (387, 388)				201001
195	13518389 (389, 390)	Novel Protein eim Chank			Z045U3
}	(200 200 200 200 200 200 200 200 200 200	1914980892[gb]AAD35474.1[AE00171 - (AE001718) ABC		transport	264636
1					
<u> </u>	83003368 (381, 392)	Novel Protein sim. GBank gi[1705461 sp P53656 BIOA_ERWHE - ADENOSYLMETHIONINE-8-AMINO-7-OXONONANOATE	Contains protein domain (PF00202) - gaba Aminotransferases dass-III pyridoxal obosobate	gaba	264600, 264689, 264638
		AMINOTRANSFERASE (7.8-DIAMINO-PELARGONIC ACID AMINOTRANSFERASE) (DAPA AMINOTRANSFERASE)			
197	80248665 (393, 394)	Novel Protein sim. GBank	Contains protein domain (PF00365) - kinase	kinase	264602, 264682, 264692, 18108374
		gij3122305jspjQ27778jK6PF_SCHMA - 6- PHOSPHOFRUCTOKINASE (PHOSPHOFRUCTOKINASE) (PHOSPHOHEXOKINASE)	Phosphofructokinase		
198	79163635 (395, 396)				264636
199	78890715 (397, 398)	Novel Protein sim. GBank gil1781203 emb CAB06110  -	Contains protein domain (PF00393) -		265008
		(Z83859) gnd [Mycobacterium tuberculosis]	6-phosphogluconate dehydrogenases		
20	79413849 (399, 400)	Novel Protein sim. GBank gij2642222 (AF030885) - telomere-associated recQ-like helicase [Ustilago maydis]		UNCLASSIFIED	264595, 264596
201	86945924 (401, 402)			UNCLASSIFIED	29331826 265007 264512 33657402
		(Y14573) ring finger protein (Hordeum vulgare)			264596, 265017, 18108351, 264682, 264683,
					284767, 264629, 55810764, 284634, 264635,
İ					58182323, 60432113, 22279000

550	705 CALL 2402 4041	Minist Destrict of the Only				
: 1			Contains protein domain (PF01644) - synthase Chilin synthase	synthase	264600	
203	79843927 (405, 406)	Novel Protein sim. GBank gij1504042 dbj BAA13220  - (D86984) similar to yeast adenylate cyclase (S56776)			22278995, 29331822, 29331825, 29331827, 264906, 21908754, 264883, 21908768,	,
204	79855186 (407, 408)	213,482		INC. ACCIEIED	21906769, 35696423, 264556	
22	10090583 (409, 410)	Novel Protein sim. GBank gil2833808 emb CAB13310  - (Z99111) similar to hypothetical proteins (Bacilus subilis)		transport	264909	
506	8758473 (411, 412)			Г	254504	_
7.	20754522 (413, 414)	Novel Protein sim. GBank gi 2134381 pir  S60678 -		UNCLASSIFIED	264556	
208	20289261 (415, 416)				264605	
99	80071069 (417, 418)	Novel Protein sim. GBank			264606 264600	
		gi 2501040jsp O05814 SYP_MYCTU - PROLYL-TRNA SYNTHETASE (PROLINETRNA LIGASE) (PRORS)			204003, 204003	
210	80168800 (419, 420)				264905, 264907, 264909, 264766, 264687,	
	, , , , , , , , , , , , , , , , , , , ,				264691, 264629, 18108374, 264638	
	60034539 (421, 422)				263978	_
2	82442474 (423, 424)	Novel Protein sim. GBank		UNCLASSIFIED	264508 264905 264906 264907 26490R	_
		gi 5031809 ref NP_005538.1 plSLR - immunoglobulin superfamily containing leucine-rich repeat			264600, 284762, 264534, 264632, 264634, 264635, 264634, 264636, 264636, 264688	
213	80249562 (425, 426)	Novel Protein sim, GBank	Contains protein domain (PE00130)	icomoraco	27278006 264608 264600 264602 264602	_
	TON FOULTOON	gij3122359jspj033123 LEU2_MYCLE - 3. ISOPROPYLMALATE DEHYDRATASE LARGE SUBUNIT (ISOPROPYLMALATE ISOMERASE) (ALPHA-IPM ISOMERASE) (IPMI)	Aconiase family (aconitate hydratase)	Somerase	224.03990, 204300, 204600, 204603, 284603, 284605, 33657023, 264565, 264486	
4	80079381 (427, 428)	Novel Protein sim, GBank	Contains protein domain (PF00118) - leph	qaa	264600 264693	_
	·	gi[116236 sp P19421 CH60_COXBU - 60 KD CHAPERONIN (PROTEIN CPN60) (GROEL PROTEIN) [(HEAT SHOCK PROTEIN B)	TCP-1/cpn60 chaperonin family			
212	14973283 (429, 430)			UNCLASSIFIED	264629	
912	80177716 (431, 432)	Novel Protein sim. GBank gi 3417297 (AC002310) - Unknown gene product [Homo sapiens]	Contains protein domain (PF00096) - dna_ma_bind Zinc finger, C2H2 type	dna_ma_bind	264448	
217	78603634 (433, 434)	Novel Protein sim. GBank gi 2506924 sp P49754 VP41_HUMAN - VACUOLAR ASSEMBLY PROTEIN VPS41 HOMOLOG (S53)			264508	
218	80258475 (435, 436)	Novel Protein sim. GBank gi[1173286]splP38106 RSEA_ECOLI - SIGMA-E FACTOR NEGATIVE REGULATORY PROTEIN		rnapolymerase	264594	
218	20438797 (437, 438)	Novel Protein sim. GBank gij1781097jembjCAB06231j - (283864) gitB [Mycobacterium tuberculosis]		synthase	264604	
022	13499572 (439, 440)	Novel Protein sim. GBank gij2984703 (AF052427) - unknown [Trypanosoma cruzi]		nudeaseinhib	264689	
	11287498 (441, 442)	Novel Protein sim. GBank gil4587313 db  BAA76709.1  -  (AB025248) alpha-1,2-mannosidase [Bacilius sp. M-90]	-	UNCLASSIFIED	264555	

247	79873185 (493, 494)	Novel Protein sim. GBank gij 1839006jemb CAB06648j - (285982) argB (Mycobacterium tubercutosis)		kinase	264909, 264691, 35696423, 18108387
248		Novel Protein sim. GBank gij1168574 sp P42464 ATPB_CORGL - ATP SYNTHASE BETA CHAIN		synthase	35596286, 264907, 264511, 264602, 264768, 264688, 265021, 35695855, 18108385
248	79764645 (497, 498)			UNCLASSIFIED	264907, 264910, 265011, 264762, 264636
250	79519980 (499, 500)				21906768, 264692
251	84359489 (501, 502)			UNCLASSIFIED	52645156, 29331822, 29331824, 52644045, 265018, 21906765, 21906768, 265020, 27486261, 27486265, 35695763, 18108376, 264556, 264559, 284565
252	79737756 (503, 504)	Novel Protein sim. GBank gij3327166 dbj BAA31651  - (AB014576) KIAA0676 protein [Homo sapiens]			264685, 264687, 264632
253	20443124 (505, 506)	Novel Protein sim. GBank gi 3036880 emb CAA18513  - (AL022374) putative ATP-dependent DNA helicase  Streptomyces coelicolor		helicase	264604
254	80027421 (507, 508)	Novel Protein sim. GBank gij3915488 spj034981 YJMB_BACSU - HYPOTHETICAL SYMPORTER IN COTT-RAPA INTERGENIC REGION		UNCLASSIFIED	264508, 264906, 264602, 264687, 265021, 264486
255	11398315 (509, 510)	Novel Protein sim. GBank gil1665720 dbj BAA04134  - (D17312) diarrheal toxin (Bacillus cereus)		UNCLASSIFIED	264593
256	80028158 (511, 512)	Novel Protein sim. GBank gil465787 sp P34422 YL31_CAEEL - HYPOTHETICAL 86.0 Prolyl oligopeptidase family KD PROTEIN F4489.1 IN CHROMOSOME III	Contains protein domain (PF00326) - peptidase Protyl oligopeptidase family	peptidase	264602, 264592
257	20289282 (513, 514)	Novel Protein sim. GBank gi[1172039]spiP42315]SCOA_BACSU - PROBABLE SUCCINYL-COA:3-KETOACID-COENZYME A TRANSFERASE SUBUNIT A (SUCCINYL COA:3-OXOACID COA-TRANSFERASE) (OXCT A)	Contains protein domain (PF01144) - transferase Coenzyme A transferase	transferase	264605
258	20459464 (515, 516)	Novel Protein sim. GBank giļ3127836 emb CAA18902  - [AL023496] hypothetical protein (Streptomyces coelicolor]		UNCLASSIFIED	264604
529	79910152 (517, 518)			collagen	264681, 264686, 264692
	20208437 (519, 520)			UNCLASSIFIED	264692, 264558
	20265063 (521, 522)	Novel Protein sim. GBank gif1237611sp[P24221 HUTH_STRGR - HISTIDINE AMMONIA-LYASE (HISTIDASE)	Contains protein domain (PF00221) - UNCLASSIFIED Phenylalanine and histidine ammonia Mases	UNCLASSIFIED	264600
262	80189317 (523, 524)			UNCLASSIFIED	265017, 264369
	88095045 (525, 528)	Novel Protein sim. GBank gil3924708 emb CAA84646 . (235597) Weak similarity with sea squirt nidogen precursor protein (blasto score 71); cDNA EST EMBI -102069 comes		UNCLASSIFIED	26448B, 264905, 264806, 264807, 264908, 284909, 264512, 264910, 264758, 264596, 264604, 264758, 264584, 264758, 264767
		from this gene; cDNA EST EMBL:D76135 comes from this gene; cDNA EST EMBL:D73147 comes from this gene; cDNA EST EMB			264763, 264764, 264288, 264769, 264768, 264769, 2646764, 264689, 264693, 264628, 264769, 2646999, 264699, 264699, 264699, 264699, 264699, 264699, 264699, 264699, 264699, 264699, 264699, 264699, 264699, 264699, 264699, 2646999, 2646990, 2646990, 2646990, 2646990, 2646990, 2646900, 26469000000000000000000000000
					204034, 204033, 204333, 204030, 204038, 264639
¥	87370826 (527, 528)	Novel Protein sim. GBank gil3043734 dbj BAA25531  -  (AB011177) KIAA0605 protein [Homo sapiens]	Contains protein domain (PF00047) - protease Immunoglobulin domain	protease	264259, 264908, 21906754, 265018, 265019, 265020,

				264400 2000000 20124824 56182181
592	95355646 (529, 530)	Novel Protein sim. GBank gil4589624 dbj BAA76834.1  -   (AB023207) KIAA0990 orotein [Homo sapiens]	Kinase	35696052, 264508, 264905, 264908, 264907,
			Ξ.	66712502, 264908, 264909, 264511, 264512.
		,		264910, 264592, 264595, 264758, 264596.
				55811386, 264600, 265017, 264603, 264604.
			•	264605, 264760, 18108351, 264762, 264681.
		•		264764, 264288, 284766, 264768, 264769.
_				21906765, 21906767, 21906769, 265020,
	•			264691, 33657023, 33657108, 33657182.
				264628, 35696423, 35695855, 264630,
				284631, 284632, 264634, 264635, 284636,
				264666 284838 83373044 58528488
				COLORS CONTROL DEARER DRAARE
				67 100310, 204304, 204300, 204400
266	79588075 (531, 532)			2646UU
287	1136222 (533 534)			264828
388	70000568 (535, 536)			264687, 264769, 264689
	00036040 (537 538)		UNCLASSIFIED	264602
Ĉ,	60023610 (331, 339)	11.00	INCI ASSIFIED	264693
2	84361144 (539, 540)	Nover Protein sim. Gbank  gi 4507367[ref]NP_003182.1 pTARS - threonyl-tRNA		
٦	78552301 (541, 542)		UNCLASSIFIED	264909, 264693
373	0874778 /543 544)	Nowal Protein sim GBank	synthase	264908
į	(100)0111100	ajl4980738jabjAAD35331,1JAE00170 - (AE001707) glucose-		
273	12840694 (545, 546)	Novel Protein sim. GBank	UNCLASSIFIED	264688
		gil1168224[sp]P44569[5NTD_HAEIN - PROBABLE 5:		
	1073 2737 07070300	NOCKEO HONSE TRECORSON		264564
2)7	39324246 (347, 346)		INCI ASSIFIED	264907 264908 264909 264766 264768.
72	82787041 (549, 550)	Novel Protein sim. GBank gij3253139 (AF 005359) - Iranslation initiation factor efF2C [Oryctolagus cuniculus]		264691, 264632, 264636
276	86671073 (551 552)	Novel Protein sim. GBank		265008, 60432229
2		gil134920 sp P21997 SSGP_VOLCA - SULFATED		
		SURFACE GLYCOPROTEIN 183 (SSG 183)	ribosomalorot	264600 18108387
277	80078735 (553, 554)	Novel Protein sim, Gbank gij128021[sp]P20964[OBG_BACSU - SPO08-ASSOCIATED		
	1023	GTP-BINDING PROTEIN	INCI ASSIFIED	264689
8	12866947 (333, 330)		o-release	284578 264604 21906764 264638 264557.
279	95292719 (557, 558)	Novel Protein sim. GBank gif79839[pir] S03812 - uvrb	noceaso	264404
18	(003 033) \$10000			284259
	2003017 (339, 300)	Norm Dratein eim CBank		18108392, 264634, 264555, 264556, 264557.
107	00248388 (301, 302)	NOVEL TIOUSING SAID. COOKING		264558
		1943123190/jspju 19394j1 LNZ_CAEEL - HTTOTAETICAL 148.2 KD TRP-ASP REPEATS CONTAINING PROTEIN		
		D2013.2 IN CHROMOSOME II		
282	18598682 (563, 564)		UNCLASSIFIED	265019
283	20614211 (565, 568)		UNCLASSIFIED	264555

284	91212160 (567, 568)	Novel Protein sim. GBank gil2429094 (U58632) - acetyl	Contains protein domain (PF00300) - LUNCI ASSIFIED	Γ	35696052 29331828 264508 264905
		xylan esterase; AxeA [Thermotoga neapolitana]	Phosphoglycerate mutase family		264600, 264602, 264605, 284682, 264764
					56181562, 21906764, 18108376, 264636.
7					264559, 18108387
285	8757940 (569, 570)			UNCLASSIFIED	264603
	80503235 (571, 572)	Novel Protein sim. GBank gi 2072674 emb CAB08305  - (295120) thiE [Mycobactertum tuberculosis]	Contains protein domain (PF00270) - ATPase_associated 35696052, 264769, 264638 DEAD/DEAH box helicase	ATPase_associated	35696052, 264769, 264638
287	12745521 (573, 574)			UNCLASSIFIED	264689
	20756502 (575, 576)	Novel Protein sim. GBank gil765323 bbs 157676 - (S74439)		l	264557
		silk fibroin heavy chain {C-terminal} [Bombyx mori=siltworms, Peptide Partial, 633 aal [Bombyx mori]			
289	80043804 (577, 578)	Novel Protein sim. GBank gi[1870009 emb CAB06860] -	Contains protein domain (PF00440) - ribosomalprot	ribosomalprot	264593, 264600
		(toberculosis)	bacterial regulatory proteins, teta family		
290	80430175 (578, 580)			UNCLASSIFIED	264768
	20747431 (581, 582)	Novel Protein sim. GBank		UNCLASSIFIED	264601
		gij2506664 spiP40120jYDCG_ECOLI - 59.4 PROTEIN IN TRG-RIMI INTERGENIC REGION PRECI IRSOR			
282	80052555 (583, 584)	Novel Protein sim. GBank gil625182 (L39015) -		UNCLASSIFIED	284605
		mitochondrial glutamyl-tRNA synthetase [Saccharomyces			
		cerevisiae]			
293	80062519 (585, 586)	Novel Protein sim. GBank		helicase	264909, 264605, 264687, 264689, 264692
		gil1718065[sp]P53528JUVRD_MYCLE - PUTATIVE DNA			
284	79830303 (587 588)	Novel Protein elm Chank	Cartaina activity domain (BE00009)		3000000 100000 000000
		gil17222Sp10040[CRB_DROME - CRUMBS PROTEIN PRECIDE OF ORES	EGF-like domain	allahollo	55811576
282	79444180 (589, 590)	Bank			52644507, 29331822, 264592, 265020,
ļ	1002 7047 0505005	(D82384) a variant of TSC-22 [Gallus gallus]	4		264639
987	79607076 (591, 592)	Novel Protein sim. GBank gi 3649789 db  BAA33403  -  (AB012226) SecA (Vibrio alginolylicus		synthase	264508
297	79631297 (593, 594)	Novel Protein sim. GBank gil5689967 emb CAB52004.1 -		UNCLASSIFIED	264905, 264687, 264638
		(ALTUSGOS) putative membrane protein (Streptomyces coelicotor A3(2))			
298	80418898 (595, 596)			UNCLASSIFIED	264905, 264691, 264639, 264766

299		Novel Protein sim. GBank gij220637(dbj BAA01477  - (D10627) zinc finger protein [Mus musculus]	Contains protein domain (PF00096) -		264488, 263994, 56994075, 22278997, 22278998, 22278999, 20281099, 29331824, 29331826, 29331826, 29331826, 29331826, 29331826, 29331826, 29331826, 264905, 264907, 264908, 264595, 264907, 264908, 264595, 264596, 264764, 265008, 264764, 264768, 264768, 264768, 264769, 264768, 264768, 264768, 264769, 264768, 264687, 3618182, 264769, 3657109, 27486261, 18108370, 264628, 264634, 2646444, 2646444, 2646444, 2646444, 2646444, 2646444, 2646444, 2646444, 2646444, 2646444, 2646444, 264
9 9 9	20711340 (599, 600)			UNCLASSIFIED	264602
301	13511332 (601, 602)	Novel Protein sim. GBank gill 145922 (M20981) - Iron dicitrate transport protein precursor [Escherichia coil]		transport	264687
302	9875260 (603, 604)	Novel Protein sim. GBank gif1174661[spiP44594 TGT_HAEIN • QUEUINE TRNA• RIBOSYLTRANSFERASE (TRNA•GUANINE TRANSGLYCOSYLASE) (GUANINE INSERTION ENZYME)			264908
33	79574895 (605, 606)				264689
ğ g	20711344 (607, 608)	Novel Protein sim. GBank gij67985 pirj HJNVAV - helicase (EC 3.6.1) - Autographa californica nuclear polyhedrosis virus		helicase	264602
g g	80412520 (609, 610)	Novel Protein sim. GBank gij728867[sp P40602 APG_ARATH - ANTER-SPECIFIC PROLINE-RICH PROTEIN APG PRECURSOR			264763
<u>g</u>	8515876 (611, 612)	Novel Protein sim. GBank gij1657554jgbjAAB18082.1 - (U73857) hypothetical protein [Escherichia coli]	-	UNCLASSIFIED	263978
200	80222901 (613, 614)			UNCLASSIFIED	265010, 21906768, 265020, 18108374, 263977
908	80064305 (615, 616)	Novel Protein sim, GBank gi 1710612 sp 010793 RNH2_MYCTU • PROBABLE RIBONUCLEASE HII (RNASE HII)	Contains protein domain (PF01351) - nuclease Ribonuclease HII	nuclease	264910, 264600, 264605, 264687, 264689, 264638, 18108387
308	80504136 (617, 618)	Novel Protein sim. GBank gil5420387 emb[CAB46679.1] - (AJ243459) proteophosphoglycan [Leishmania major]			264769
9	80053616 (619, 620)				264603
115	11090659 (621, 622)	Novel Protein sim. GBank gi 1144522 (U34957) - phosphoribosylaminolmidazolesuccinocarboxamide <u>synithase [Mycobacterium tuberculosis]</u>		synthase	264602
315	80054347 (623, 624)			UNCLASSIFIED	264566
3	(979, 679) (979) (979)				284603, 284567

-			Contains protein contain (Frocazo) - una_ma_omo		SCHELLS, COMPLEXOR, EGGS 1040, EGGS 2003.
_			CATA ZIIIC III gal		60432229, 33657402, 60433438, 33109954.
					265011, 265017, 264603, 265018, 264288,
_					264766, 264692, 35695763, 264628, 264829,
					264639, 60170394, 22279002, 264568
186	82356091 (629, 630)	Novel Protein sim. GBank gil 1652620 dbj BAA17540  -			264508, 264600, 264762, 264687, 264768,
		(D90907) pyridine nucteotide transhydrogenase beta			52644229, 284769, 264689, 264635, 264636,
-		subunit [Synechocystis sp.]			264638, 264486
7	78911071 (631, 632)			UNCLASSIFIED	264693
2	0466944 (633, 634)			UNCLASSIFIED	264605
		gij118244jspjP24176jDAPE_ECOLI - SUCCINYL-			
-10	04141838 (635 636)	Novel Protein sim GRank	Contains protein domain (PE00526) - Iransport	fransport	264908, 264909, 264910, 264593, 264594.
	(000,000)	110001 1 100011 5411. OBJIN 011488022010b1AAD27583 11AE11827 - (AE11827A) DNb-5	Dichostefilm (slime mold) repeats		264760, 264288, 264768, 264769, 21906769.
		[Homo sapiens]			264691, 264693, 264628, 65274791, 264635.
					264636, 264638, 83373044, 22279002,
+	17300360 (637 630)	Name   Designation of the Control of		2000000	265048
<u>-</u> _				nodenan	0.007
۴	13527575 (630 BAD)	Novel Protein eim CBank		evnihaca	764687
_	_	haven 100cm sim. Obsum hij28110331sniO053141Gl GC MYCTU - GLUCOSE-1-		Seminife	
_		PHOSPHATE ADENYLYLTRANSFERASE (ADP.	•		
		GLUCOSE SYNTHASE) (ADP-GLUCOSE			
		PYROPHOSPHORYLASE)			
۳	94134387 (641, 642)	Novel Protein sim. GBank gi[1680716 (U68234) - all-trans-		cyto450	264509, 264906, 264907, 264908, 265009.
		retinoic acid 4-hydroxylase [Danio rerio]			264596, 264764, 264628, 264634, 264635, 264638, 264630, 83373304, 264567
+	36400000 (040 044)	00 Old 1 (93050) 1 33000 this stock of the Charles		COLDINA POPICION	ECO111EN 264601 60421428 55810764
	qo489053 (643, 644)	Novel Protein sim. GBank gij1160355 (U33056) - UNC-89  [Caenorhabditis elegans]		UNCLASSIFIED	33611130, 264631, 60431326, 33610764
۳	94653725 (645, 646)			UNCLASSIFIED	264488, 265009, 264593, 264628, 264635
H	79174383 (647, 648)				264687
	79862691 (649, 650)			UNCLASSIFIED	264693
ř	28774974 (651, 652)			UNCLASSIFIED	264288, 18108385
<del></del>	79776267 (653, 654)	Novel Protein sim. GBank gij451544 (U04267) - proline-rich cell wall protein (Gossyplum barbadense)			284488, 264905, 264509, 264910
<del>ا</del> ≃	80253202 (655, 656)			UNCLASSIFIED	264592
Ť	10173821 (657, 658)			UNCLASSIFIED	284510
<del>                                     </del>	88597767 (659, 660)	Novel Protein sim. GBank gil4191358 (AF087825) - claudin-		UNCLASSIFIED	264259, 264908
۳	7076/000 /664 663)	March Order of Contract Contra		to to the contract of the cont	SEADIO SEAEBY SEAEBO SEAESE SEAEET
-	19134666 (601, 602)	Nover Protein Sim. Chank gilou/41 priglocus 1.2 - regulatory protein whiB - Streptomyces coelicolor		uanscripuacio	204910, 204001, 204009, 204000, 204001
ŕ	80071440 (663, 664)	Novel Protein sim. GBank		reductase	35696423, 264636, 264638, 264565
		BIJ114049 sp P19480 AHPF_SALTY - ALKYL			
		HYDROPEROXIDE REDUCTASE SUBUNIT F (ALKYL HYDROPEROXIDE REDUCTASE F524 PROTEIN)			
۲	13009555 (865, 666)				264687

264905, 284600, 264604, 264486	264907, 264603, 264693, 18108374, 264636, 18108387	265009, 264766, 264686	264602	264569, 18108397, 22278998, 29331822, 20281099, 29331824, 58182181, 68714117, 29331825, 35686052, 28331828, 264508, 264906, 264906, 264906, 264906, 264906, 265008, 264909, 265008, 26400, 264501, 264681, 264764, 265018, 265019, 264682, 265017, 264682, 264682, 264681, 264762, 18108351, 264682, 264682, 264689, 264689, 264686, 264686, 264637, 264632, 264638, 2	264593	264905, 265019, 264769, 18108374		264605	264603, 264604	264605		264593	264905
dehydrogenase	esterase	UNCLASSIFIED		nud_recpt	ydə	ubiquitin		Isomerase			UNCLASSIFIED	histone	dehydrogenase
1022228 pir  532227 - glutamate   Contains protein domain (PF00208) - dehydrogenase   C 1.4.1.4) - Corynebacterium   Glutamate/Leucine/Phenylalanine/Va   line dehydrogenase				Contains protein domain (PF00249) - Inucl_recpt Myb-like DNA-binding domain				Contains protein domain (PF00290) - Isomerase Tryptophan synthase alpha chain					Contains protein domain (PF00208) - dehydrogenase Glutamate/Leucine/Phenylalanine/Valline dehydrogenase
Novel Protein sim. GBank gij322228 pir  S32227 - glutamate  dehydrogenase (NADP+) (EC 1.4.1.4) - Corynebacterium glutamicum	Novel Protein sim, GBank gijz193938jembjCAB09602j - (296800) gipQ2 [Mycobacterium tuberculosis]			Novel Protein sim. GBank gi[5454074 ret NP_006303.1 pSMRT - silencing mediator for Myb-like DNA-binding domain retinoid and thyroid hormone receptors	Novel Protein sim. GBank gil4001713 dbj BAA35087.1  - [AB015879] DnaK [Porphyromonas gingivalis]		gi[2842699]sp[092353]UBPC_SCHPO - PUTATIVE UBIQUITIN CARBOXYL-TERMINAL HYDROLASE C6G9.08 (UBIQUITIN THIOLESTERASE) (UBIQUITIN-SPECIFIC PROCESSING PROTEASE) (DEUBIQUITINATING ENZYME)	Novel Protein sim. GBank gil2688580 (AE001166) - conserved hypothetical protein [Borrella burgdorferi]	Novel Protein sim. GBank gil 1684738jembjCAA70601   - (Y09452) Yed I hypothetical protein [Pseudomonas syringae]		Novel Protein sim. GBank gij2117275 embjCAB09104  - (295818) hypothetical protein Rv0807 [Mycobacterlum tuberculosis]	Novel Protein sim. GBank gij3023317[spjQ48935[aPHA_MYCRA - ACETYLPOLYAMINE AMINOHYDROLASE	Novel Protein sim. GBank gil4239787 emb CAA75437  - (Y15166) NADP-glutamate dehydrogenase [Pseudomonas aemginosa]
	80057028 (669, 670)	80414319 (671, 672)	11090829 (673, 674)		11398513 (677, 678)	80504149 (679, 680)		11075198 (681, 682)	80054196 (683, 684)	20466792 (685, 686)	80428870 (887, 688)	80258853 (689, 690)	78831058 (691, 692)
		Т	Г	338	338	8		74	342	343	344	345	346

			1		
	•	oxidoreductase chain 4L	DEHYDROGENASE I CHAIN K (NADH-UBIQUINONE OXIDOREDUCTASE CHAIN 11) (NIIO11)		
264769, 264602, 264604, 264508, 264762,	dehydrogenase	Contains protein domain (PF00420) - dehydrogenase	Novel Protein sim. GBank gi[2829816[sp[P95171]NUOK MYCTU - NADH	074-2222 (171, 170)	<u> </u>
264566				79750145 (725, 726)	200
	,	-	gij113764ispiP25718jAMY1_ECOLI - ALPHA-AMYLASE PRECURSOR (1,4-ALPHA-D-GLUCAN GLUCANOHYDROLASE)		
264688	amvlase		Novel Protein sim. GBank	13089485 (723, 724)	362
264605, 55811957, 285022			CaliNAC:polypeptide N-acetylgalactosaminyltransferase T5   [Rattus norvegicus]		
22278996, 264259, 29331822, 29331824,	transferase		Novel Protein sim. GBank gi[3510639 (AF049344) - UDP-	80584075 (721, 722)	198
264594	UNCLASSIFIED			80026748 (719, 720)	360
264604 264769	UNCLASSIFIED			80501488 (717, 718)	328
264768	UNCLASSIFIED		Novel Protein sim. GBank gi 2290990 (AF006000) - Brg1  {Bordetella pertussis}	3/032/36 (713, 716)	900
264605	oxidase		Novel Protein sim. GBank gil497637 (J03939) - cytochrome oxidase d subunit i [Escherichia coii]	60070366 (713, 714)	è
		Response regulator receiver domain	gillala/ispip18574 BVGA_BORPE - VIRULENCE FACTORS PUTATIVE POSITIVE TRANSCRIPTION REGULATOR BVGA		
264909, 264595, 264683, 22279002	transcription	Contains protein domain (PE00072) - transcriptiactor	Novel Protein sim. GBank	80043835 (711, 712)	328
284000 264505 264603	INC. ACCIETO			80046344 (709, 710)	355
264638	INCIACIEIED			56626130 (707, 708)	354
		Urease	gi 1174887lsp P42873 URE1_STAXY - UREASE ALPHA SUBUNIT (UREA AMIDOHYDROLASE)		
284804	INCIASSIFIED	Contains protein domain (PF00449) - 11NC1 ASSIETED	(AF105716) copia-type pol polyprotein [Zea mays] Novel Protein sim. GBank	80061653 (705, 706)	353
DEAEDE	protosea		Novel Protein sim. GBank gil4416302lobIAAD203071 -	11611585 (703, 704)	352
264769, 264905, 264908	glycoprotein		Nover Protein sim. GBank gilz959367 jembjCA417921 j - (AL022117) hypothetical protein [Schizosaccharomyces pombe)	(201, 101) (00)	3
			(277137) hypothetical protein Rv1277 [Mycobacterium tuberculosis]		
265009 284789 264689 18108370	nuclease		Novel Protein sim. GBank gij3261599 emb CAB00917  -	80502370 (699, 700)	320
265007				17282112 (697, 698)	349
264602, 18108351, 18108387	transport		Novel Protein sim. GBank gil1073610 pir  S47672 - ugpB protein - Escherichia coll	80020208 (695, 696)	% %
263981			gij731675 spiP38795 YHN4_YEAST - HYPOTHETICAL 80.7 KD PROTEIN IN ERG7-NMD2 INTERGENIC REGION		
265006, 265008, 265010, 265018, 263967	UNCLASSIFIED		Novel Protein sim. GBank	347 79158185 (693, 694)	Ì

365	88040288 (729, 730)	Novel Protein sim. GBank gil4929268 gblAAD33924.1 -	Contains protein domain (PF00412) -		264488, 21906766, 21906767, 55811576,
		(AF144237) LOMP protein [Homo sapiens]	LIM domain containing proteins		21906769,29148629,22278995,22278996, 264020 264022 264634 264801 264803
					33657023, 33657402, 264693, 264639,
					264594, 29331824, 264758, 18108385,
					29331827, 87168559, 265018, 22279000.
					265019, 264482, 264761, 264681, 18108351
366	81821838 (731, 732)				265017, 264757
	95357471 (733, 734)	Novel Protein sim. GBank	Contains protein domain (PF01602) - glycoprotein		60424179, 65274572, 56182575, 22278994.
		gi 4503843 ref NP_003908.1 pG2AD - UNKNOWN	Adaptin N terminal region		56994075, 22278998, 264259, 29331822,
					29331824, 56182181, 60424269, 66714117.
				-	29331825, 60432289, 29331826, 29331827,
					29331828, 264905, 264828, 56182435,
					265006, 264512, 265008, 264591, 55812038,
					55811386, 265010, 87168559, 265017,
					265018, 264604, 265019, 55811150, 264448,
		-			264369, 264288, 264686, 264768, 56181562,
					21906768, 21906769, 55811957, 35695917,
				. <del></del>	265022, 60170615, 33657023, 65274620.
					18108365, 263967, 33657109, 33657349,
					35695763, 264628, 18108376, 55811576,
					65274791, 35695855, 56182323, 83373044.
					60432113 264563 264564 264567
368	79607265 (735, 736)				264509
200	1952 7777 7381	Novel Protein sim CBank		INCI ASSIFIED	264508 264604 264605 264636
}	(201.101)	gij3913029jspjP94967jALR MYCSM - ALANINE			
		RACEMASE			
370	88090966 (739, 740)	Novel Protein sim. GBank gi 3249559 (AF018261) - EH			264905, 264592, 264605, 264766, 264691
	•	domain binding protein Epsin [Rattus norvegicus]			
371	95292599 (741, 742)	Novel Protein sim. GBank gil2995299jemb[CAA18328] -	Contains protein domain (PF01715) - Iransferase	transferase	264905, 264906, 264510, 264600, 264601,
		(AL022268) putative IRNA delta(2)-	IPP transferase		264602, 264603, 265018, 264604, 264605,
		Isopentenylpyrophosphate transferase (Streptomyces			265021, 264692, 264636, 264564
		coelicolor			
372	80021107 (743, 744)		_		264564
		(gi 2506393 sp P31576 FIXX_ECOLI - FERREDOXIN LIKE   PROTEIN	•		
373	79863766 (745, 746)			UNCLASSIFIED	264909
374	79847568 (747, 748)	Novel Protein sim. GBank gij3341640jembjCAA13164j -		UNCLASSIFIED	264905, 264906
		(AJ231122) z61f [Vibrio cholerae]			
375	91230181 (749, 750)	Novel Protein sim. GBank gi 5456934 gb AAD43716.1 -		cadherin	65274572, 264259, 29331826, 56182435.
		(AF152322) protocadherin gamma A2 [Homo sapiens]			60433356, 60433438, 264757, 55812038, 264758, 55811957, 264690, 33657023
378	80505214 (751, 752)	Novel Protein sim. GBank gil1805408 dbi BAA08970  -		UNCLASSIFIED	264769
		(D50453) homologues to nitrile hydratase region 3'-			
		hypothetical protein P47K of P. chlororaphis [Bacillus subtilis]			
377	10339083 (753, 754)				264906

							264595, 265017, 265021, 264638, 87168518. 22279002	264259, 29331822, 60432289, 29331827,	264266, 264766, 263867, 63274781, 35695855, 263981, 83373044, 264567				35696286, 264905, 66712502, 60432229,	264593, 60433356, 264686, 264688, 21906765, 264891, 22279000, 264482	996, 264630, 264556,				4				265007, 265009, 264508, 264556, 264629, 264766	18108398, 22278995, 22278996, 56994075, 22278999, 264259, 29331824, 29331826, 264905, 264908, 265007, 265009, 265009, 21906754, 33557084, 265017, 264448, 264288, 264766, 21906765, 21906765, 21906765, 21906767, 265021, 33657023, 3460423, 34604655
	265008, 284555	264769		264604	264684	264592	264595, 265017, 2 22279002	264259, 29331822	35695855, 263981	264692	264906	264760	35696286, 26490	264593, 6043335(21906765, 26469	18108394, 22278 22279002	264600	264482	264908	264602, 21906764	264693	264508, 264563	264556	265007, 265009, 264766	18108398, 22278 22278999, 26426 264905, 264908, 21906754, 33557 264288, 264768, 21906767, 26502
	UNCLASSIFIED	synthase		dehydrogenase	UNCLASSIFIED			UNCLASSIFIED			dehydrogenase	UNCLASSIFIED	нотеорох		UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	synthase		UNCLASSIFIED	struct		Iransport
		Contains protein domain (PF00289) - synthase Carbamoyl-phosphate synthase	CPSase)	Contains protein domain (PF01011) - dehydrogenase PQQ enzyme repeat							Contains protein domain (PF00465) - Iron-containing alcohol dehydrogenases	Contains protein domain (PF01841) - UNCLASSIFIED Transglutaminase-like superfamily	Contains protein domain (PF00646) - homeobox	F-box domain.								Contains protein domain (PF00047) - struct		Contains protein domain (PF00153) - Iransport Mitochondrial carrier proteins
		Novel Protein sim. GBank gi 1076013 prr  A49930 - carB protein homolog - Mycobacterium bovis (strain BCG)		gij216556jdbj BAA02174  - genase [Escherichia coli]			Novel Protein sim. GBank gij3327136jdbjjBAA31636j - (AB014561) KIAA0681 protein [Homo sapiens]				Novel Protein sim. GBank gij1073456 pir  S47810 - probable   Contains protein domain (PF00465) - dehydrogenase alcohol dehydrogenase (EC 1.1.1.1) - Escherichia coli dehydrogenases	gi 1460074 emb CAB01049  - ein Rv2566 [Mycobacterlum	Novel Protein sim. GBank gil4240169idbilBAA74863.11 -	itein [Homo sapiens]			Novel Protein sim. GBank gil854065 emb CAA58337  - (X83413) U88 [Human herpesvirus 6]		Novel Protein sim. GBank gij3378523[emb CAA08867] - (AJ009832) cyclomaltodextrinase glucanotransferase [Thermotoga neapolitana]			Novel Protein sim. GBank gi 2677780 (U70327) - unknown [Paretroplus polyactis]	Novef Protein slm. GBank gi4507909 ref NP_000368.1 pWAS  - Wiskott-Aldrich syndrome (ecezema-thrombocytopenia)	Novel Prolein sim. GBank gij1518458 (U45998) - mitochondrial solute carrier [Onchocerca volvulus]
	٦	80503437 (757, 758)		80060937 (759, 760)	11769027 (761, 762)	80054377 (763, 764)	83259025 (765, 766)	95314255 (767, 768)		10237679 (769, 770)	79633434 (771, 772)	17960637 (773, 774)	87741378 (775, 778)		79316971 (777, 778)	80079949 (779, 780)	7657302 (781, 782)	79796056 (783, 784)	33206031 (785, 786)	10104463 (787, 788)	80229010 (789, 790)	20436224 (791, 792)	80417014 (793, 794)	91230517 (795, 796)
ſ	378 8			380	381	Γ		384		385		387	388		388	Т			393	Т	385		397	398

Novel Protein sim. GBank gij335091[dbj BAA31995] - Contains protein domain (PF00370) - (AB028939) KiAA1075 protein [Homo saplens] Novel Protein sim. GBank gij28935[sp]74 aurily of carbohydrate kinases Contains protein domain (PF00360) - (AL0091999) KiAA1075 protein [Homo saplens] Sic homology domain 2 (AL0091999) draE2 [Mycobecderium tuberculosis] Contains protein domain (PF0017) - Sic homology domain 2 (AL0091999) draE2 [Mycobecderium tuberculosis] Contains protein domain (PF00159) - Pancreatic homone peptides	284592, 264595	18108394, 56182575, 22278995, 22278997, 22278999, 264289, 265006, 265007, 265009, 60432229, 33657402, 21908754, 265010, 265017, 265018, 265019, 18108351, 18108357, 21906765, 265021, 265022, 264692, 33657023, 18108370, 65274791, 264634, 264636, 265027, 264636, 265027, 264636, 265027, 264636, 265027, 264636, 265027, 264636, 265027, 264636, 264646, 264646, 264646, 264646, 264646, 264646, 264646, 264646, 264646, 264646, 264646, 264646, 264646, 26	264594	65274572, 56182575, 35696286, 22278996, 22278998, 264093, 264059, 29331822, 29331824, 29331826, 29331826, 29331827, 29331827, 29331827, 29331828, 264906, 264907, 264909, 265006, 264511, 265007, 265008, 264910, 264591, 265010, 265011, 87168559, 265017, 265018, 265019, 264681, 18108351, 264682, 264448, 264288, 264681, 18108351, 264682, 264448, 264288, 264684, 264766, 264767, 264688, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906767, 21906765, 21906765, 21906767, 21906765, 21906765, 21906767, 21906765, 21906765, 21906767, 21906765, 21906765, 21906767, 21906765, 21906765, 21906767, 21906765, 21906765, 21906765, 21906765, 21906767, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 244633, 264539, 18108385, 264539, 18108385, 264538, 264539, 18108385, 264538, 264539, 18108385, 264538, 264546, 244655, 264546, 2446455, 244565, 2	264768, 264632, 264639, 264563	265009, 264682	18108357, 264693	264769	264600	264259
Novel Protein sim. GBank gij338091jdbjjBAA31995j - (AB015974) giycerol kinase [Pseudomonas tolaasii] Novel Protein sim. GBank gij728935[spjP39192]ALU5_HUMAN - IIII ALU SUBFAMILY SC WARNING ENTRY IIII Novel Protein sim. GBank gij4928292[gbjAA033527.1]AF13211 - (AF132117) FhuA [Staphylococcus aureus] Novel Protein sim. GBank gij569487[dbjjBAA83027.1] - (AB028998) KIAA1075 protein [Homo sapiens]  Novel Protein sim. GBank gij2681649[embjCAA15755] - (AL009198) dnaE2 [Mycobacterium tuberculosis]	kinase	cadherin	transport	phosphalase	UNCLASSIFIED	polymerase	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	helicase
Novel Protein sim. GBank gil (AB015974) glycerol kinase Novel Protein sim. GBank gil 728835[sp]P39192]ALU5 SC WARNING ENTRY !!! [Staphylococcus aureus] Novel Protein sim. GBank gil 49282999] KIAA1075 protein sim. GBank gil 4928999] KIAA1075 protein sim. GBank gil 49289999] Novel Protein sim. GBank gil 48028999] Mare GBank gil 48009199] dna E2 [Mycoba	Contains protein domain (PF00370) - FGGY family of carbohydrate kinases	Contains protein domain (PF00560) - Leucine Rich Repeat		Contains protein domain (PF00017) - Src homology domain 2			Contains protein domain (PF00159) - Pancreatic hormone peptides			
1 (801, 802) 1 (801, 802) 1 (801, 802) 1 (803, 804) 1 (803, 804) 1 (803, 805) 1 (809, 810) 1 (809, 810) 1 (811, 812) 1 (813, 814) 1 (813, 814)	Novel Protein sim. GBank gij3358091/dbj BAA31995  - (AB015974) glycerol kinase [Pseudomonas tolaasii]	Novel Protein sim. GBank gi 728835 sp P39192 ALU5_HUMAN - !!!! ALU SUBFAMILY SC WARNING ENTRY !!!!	Novel Protein sim. GBank gli4928292igbtAAD33527.1JAF13211 - (AF132117) FhuA [Staphylococcus aureus]	Novel Protein słm. GBank gij5689487 dbj BAA83027.1  -		Novel Protein sim. GBank gi 2661649 emb CAA15755  - (AL009198) dnaE2 [Mycobacterium tuberculosis]				Nover Protein Sim. GBank gij3047117 (AF058919) - similar to ATP-dependent RNA helicases [Arabidopsis thaliana]
400 94117499 400 94117499 401 1139749 402 9542029- 402 8643991- 403 8043991- 405 7947128 406 78634177- 407 80478221 407 80478221	80055278 (797, 798)	94117490 (789, 800)	11397491 (801, 802)	95420294 (803, 804)	44809843 (805, 805)	79471280 (809, 810)	79634172 (811, 812)	80478229 (813, 814)	5840527 (817, 816)	

95357496 (819, 820)	Novel Protein sim. GBank gil475016[dbj BAA06184] -	<u> </u>	UNCLASSIFIED	264489, 52646365, 52846842, 56181686,
	Isoposius sind (macining (cosso))			33838200, 32843800, 28331822, 28331824, 56182181, 29331825, 60424269, 35696052, 33656070, 264508, 264509, 264906
				264907, 264908, 52644045, 264909, 264510, 265007, 264912, 265008, 264910, 33657402,
			,	264758, 52646317, 55811388, 265010, 265011, 265017, 264604, 265018, 55811150,
				264762, 264764, 264766, 264687, 264768. 264769, 52644229, 21906766, 285020,
				2857021, 284534, 32644150, 284852, 33657023, 65274820, 33657109, 33657182,
				27486261,35695763,264628,264629, 60431528 18108376 263078 35696423
				35695855, 264632, 264634, 264635, 284637.
				264638, 264558, 264839, 56182323, 264559,
				00432113, 22273002, 204303, 204303, 264486
80501670 (821, 822)			UNCLASSIFIED	264769
80241662 (823, 824)				264907, 264910, 263973, 22279002
11076446 (825, 826)	Novel Protein sim. GBank gij3261784 embjCAB08997  -		eph	264605
82050554 (827, 828)	Novel Protein sim. GBank		dehydrogenase	18108374, 264760, 264769, 264602, 264638,
	gil129036 spiP20707 ODO1_AZOVI - 2-0XOGLUTARATE			264603, 264909, 264605
	DEHYDROGENASE E1 COMPONENT (ALPHA- KETOGLUTARATE DEHYDROGENASE)			
84453144 (829, 830)	Novel Protein sim. GBank		UNCLASSIFIED	264908, 87168518
•	gl/48683501gb/AD31273.1/AF13202 - (AF132025) rhophilin			
80402775 (831, 832)	Novel Protein sim. GBank dil2555172 (AF025543) - ArcC		kinase	264488 264600 264602 264764 264636
Ì	carbamate kinase [Rhizobium etii]			
20153797 (833, 834)		Contains protein domain (PF00145) -		264605
	gij1709171fsp[P52311[MTX2_XANOR - MODIFICATION METHYLASE XORII (CYTOSINE-SPECIFIC METHYL TRANSFERASE XORII) AM XORII)	C-5 cytosine-specific DNA methylase		
94125841 (835, 836)			UNCLASSIFIED	264689, 264693
95314273 (837, 838)			collagen	264908, 264910, 264764, 264639
37036349 (839, 840)	Novel Protein sim. GBank gij3261659[emb[CAB03751] - (ZB1368) hypothetical protein Rv2419c [Mycobacterium (tuberculosis]	Contains protein domain (PF00300) - phosphatase Phosphoglycerate mutase family	phosphatase	264769
95292942 (841, 842)	Novel Protein sim. GBank gi 2916942 emb CAA17580 -	Contains protein domain (PF00072) - phosphatase	phosphatase	264906, 264600, 264601, 264603, 264604,
	(ALDZ 1999) hypotnetical protein RV0961 (Mycobacterum tuberculosis)	Response regulator receiver domain		504160, 204109
78471293 (843, 844)		Contains protein domain (PF00118) - eph	qdə	22278996, 264682, 18108376, 18108387
	gi[231752[sp]Q00767[CH61_STRAL - 60 KD CHAPERONIN TCP-1/cpn60 chaperonin family 1 (PROTEIN CPN60 1) (GROEL PROTEIN 1) (HSP58)			
79604948 (845, 846)		П	UNCLASSIFIED	264509

<u> </u>	78966557 (847, 848)	Novel Protein sim. GBank gij4826814[ref]NP 004977.1[pXTN1 - kinectin 1 (kinesin		struct	265019
80431450 (849, 850)	. 850)	. gi 1703701 bbs 178462 - tein [rats, testes, Peptide Partial,	Contains protein domain (PF00225) - struct Kinesin motor domain		264909, 265007, 55811386, 264768, 55810764
80064522 (851, 852)	, 852)				264605, 264559
80057232 (853, 854)	3, 854)	Novel Protein sim. GBank gi[231829]sp[P29929]COBN_PSEDE - COBN PROTEIN		UNCLASSIFIED	264603, 264636
79487798 (855, 856)	, 856)			UNCLASSIFIED	264683
80091252 (857, 858)	, 858)	Novel Protein sim. GBank gi[81286 pir  S22697 - extensin - Volvox carteri (fragment)			35696423, 35695763, 35695855, 265017, 264564, 264762
80504192 (859, 860)	, 860)	Novel Protein sim. GBank gil1806154[emb]CAB06451[ -		reductase	264508 264905 264509 264908 264909
		(284395) hypothetical protein Rv0588 [Mycobacterlum tuberculosis]		•	265008, 264600, 264687, 264769, 264689, 264689, 264689
20624249 (861, 862)	1, 862)				264568
525372 (86	3, 864)				265020
81494303 (865, 866)	5, 866)	Novel Protein sim. GBank gij3123552jembjCAA18609j -		UNCLASSIFIED	264907, 264908, 264909, 264910, 264592,
		(AL022578) dJ393P12.2 (hypothelical Proline-rich protein KIAA0269 LIKE) [Homo sapiens]			264595, 264758, 264604, 264760, 264762, 264763, 264636, 264637, 22279002
94326323 (867, 868)	37, 868)		Contains protein domain (PF00169) -	Γ	55812038, 56182181, 56181562, 29331828
		gi[2495272 sp[099826 CDX2_HUMAN - HOMEOBOX PROTEIN CDX-2 (CAUDAL-TYPE HOMEOBOX PROTEIN PACES A)	PH domain		35696052, 55810764, 55811576, 65274791, 35695855, 60432113, 55811150, 264636,
A0502720 1050 0701	1070	2) (chv-3)			264766
00) 00 1300	6	NOVEL TOUGHT SHIT. OBGIN. GITTLE - L-ARABINOSE IFANSPORT SYSTEM PERMEASE PROTEIN ABAH		Iransport	264595, 264769
41085953 (871, 872)	1, 872)			INCI ASSIFIED	265020 22279002
11399291 (873, 874)	3, 874)			INCI ASSIFIED	764593
11773835 (875, 876)	5, 876)			UNCLASSIFIED	264686
80019495 (877, 878)	7, 878)				284905, 264600, 264602, 264604
79841062 (679, 880)	9, 880)	c gi 2291232 gb AAB65351.1  - ilarity to Pfam domain: PF00004 alue=3.7e-77, N=1 [Caenorhabdiits	Contains protein domain (PF00004) - ATPases associated with various cellular activities (AAA)	ATPase_associated	Contains protein domain (PF00004) - ATPase_associated  35696052, 264905, 264908, 264909, 265011, ATPases associated with various cellular activities (AAA)
20396935 (881, 882)	11, 882)	Novel Protein sim. GBank gi[5639946[gb[AAD45904.1]AF16132 - (AF161328) histidine kinase CstS (Corynebacterium diphtheriae)			264605
85281058 (883, 884)	13, 884)	Novel Protein sim. GBank gil1184790 (U46068) - von Ebner minor salivary gland protein (Mus musculus)		UNCLASSIFIED	29331830, 264909
82456427 (885, 886)	15, 886)	Novel Protein sim. GBank gi 5689893 emb CAB52056.1  - (AL109732) putative ATP-binding RNA helicase Streptomyces coelicolor A3(2)		UNCLASSIFIED	35696052, 264508, 264906, 264512, 264604, 264762, 264769, 264689, 264636
11395897 (887, 888)	37, 888)	Novel Protein sim. GBank gil1783249jdbjjBAA11726j - (D83026) homologous to citrate-sodium symport (citrate transportaret: homological (Bacilliae enhitie)		UNCLASSIFIED	264591
		ווייים בייים ווייים וייים וייים בייים ביים בייים	7		

I				ſ	20000
	79552709 (889, 890)			UNCLASSIFIED	250402
448	79810937 (891, 892)	Novel Protein sim. GBank gi[5531272]emb[CAB50897.1] - [AJ243800] WSC4 homologue [Kluyveromyces lactis]			264509
447	80438888 (893, 894)	Novel Protein sim. GBank gi 538413 (L36315) - zinc finger protein fMus musculus)	Contains protein domain (PF00096) - Iranscriptfactor Zinc finger, C2H2 type	ranscriptfactor	264768, 55811576
448	80238110 (895, 896)	Novel Protein sim. GBank gi[1542914 emb CAB02185  - 1780108) fm [Mycobacterium tuberculosis]	Contains protein domain (PF00551) - dehydrogenase Formyl transferase	dehydrogenase	264508, 264600, 264603, 264605, 264682, 264769, 18108362, 264634, 18108387
449	20460634 (897, 898)	Novel Protein sim. GBank gij118794 spiP10443 DP3A_ECOLI - DNA POLYMERASE III. ALPHA CHAIN		polymerase	264605, 264559
450	94631210 (899, 900)	Novel Protein sim. GBank gil4589508jdbjjßAA76775.1j - (AB023148) KIAA0931 protein [Homo saplens]	Contains protein domain (PF00481) - phosphatase Protein phosphatase 2C	phosphatase	65274572, 22278998, 29331824, 29331826, 264906, 264910, 264592, 52646317, 265017, 21906767, 55811957, 56528486, 22279002
451	21433609 (901, 902)			UNCLASSIFIED	264486
452	10267276 (903, 904)				264692
	52560096 (905, 906)	Novel Protein sim. GBank gil2650614 (AE001104) - conserved hypothetical protein (Archaeoglobus fulgidus)		UNCLASSIFIED	264907, 264600
2	30523022 (007 OUR)	Novel Protein cim GRank		transferase	264603
\$.	3352347 (307, 308)	NOVEL FOLEIN SIM. GBAIRA gi 2493000 sp 009450 SCOT_CAEEL - PROBABLE SUCCINYL-COA:3-KETOACID-COENZYME A TRANSFERASE PRECURSOR (3-0XOACID COA- TRANSFERASE)			
	13089692 (909 910)			UNCLASSIFIED	264687
	78563081 (911, 912)			UNCLASSIFIED	284691
457	78831273 (913, 914)	Novel Protein sim. GBank gij4468699jemb[CAB38153.1] - (AL035591) putative inlegral membrane export protein IStreptomyces coelicolori			264905
458	78581227 (915, 916)	Novel Protein sim. GBank gij3411053 (AF034863) - synaptic scaffolding molecule [Rattus norvegicus]	Contains protein domain (PF00595) - kinase PDZ domain (Also known as DHR or GLGF)	kinase	55812038, 265010, 265018, 264681
459	80567359 (917, 918)	Novel Protein sim. GBank gil4506075 ref NP_002733.1 pPRKC - protein kinase C, mu	Contains protein domain (PF00130) - kinase 3.1 pPRKC - protein kinase C, mu Phorbol esters/diacy/glycerol binding domain (C1 domain)	kinase	22278997, 264259, 29331826, 265018, 264448, 264369, 21906765, 35696423
460	79245890 (919, 920)	Novel Protein sim. GBank giji 13138jspjP25516jACO1_ECOLI - ACONITATE HYDRATASE 1 (CITRATE HYDRO-LYASE 1) (ACONITASE 1)		UNCLASSIFIED	264906
461	95287618 (921, 922)	Novel Protein sim. GBank gi 1168574 sp P42464 ATPB_CORGL - ATP SYNTHASE BETA CHAIN		synthase	264602, 264605, 264768, 264769, 265021, 33657023, 264559

462	79606589 (923, 924)	Navel Protein sim. GBank MULTIPHOSPHORYL TRANSFER PROTEIN (MTP) (CONTAINS: PHOSPHOENOLPYRUVATE-PROTEIN PHOSPHOTRANSFERASE (PHOSPHOTRANSFERASE SYSTEM. ENZYME 1); PHOSPHOCARRIER PROTEIN PROPEDIAL FRUCTOSE-SPECIFIC IIA	Contains protein domain (PF00391) - UNCLASSIFIED PEP-utilizing enzymes		264807
463	79796417 (925, 926)	Court Over II Novel Protein sim. GBank gil854065 emb CAA58337  - (X83413) U88 [Human herpesvirus 6]		UNCLASSIFIED	264905, 264906, 264908, 264910. 284591, 264595, 265011, 284632, 284635. 264636, 284637, 284638, 264639
184	182340151 (927 92R)			UNCLASSIFIED	264634
465	T	Novel Protein sim. GBank gil5689776jemb[CAB52137.1] - [A.1242832) calbain [Homo sapiens]	Contains protein domain (PF00848) - cathepsin Calpain family cysteine protease		265017, 21906764, 265020
466	20460645 (931, 932)	Novel Protein sim. GBank gil1806175[emb CAB06470] - (284395) rpsC (Mycobacterium tuberculosis)	Contains protein domain (PF00417) - inbosomatprot Ribosomal protein S3, N-terminat domain.		264605, 264559
467	80409035 (933, 934)	Novel Protein sim. GBank gij548705[spjP36949 RBSB_BACSU - D-RIBOSE-BINDING PROTEIN PRECURSOR			264764
ARR	5256220B (935 936)				264692
469	19520527 (937, 938)	Novel Protein sim. GBank gil2114024 emblCAB08957  - (295558) grcC1 (Mycobacterium tuberculosis)		UNCLASSIFIED	264488
470	80502756 (939, 940)	Novel Protein sim. GBank gij2909459jemb CAA17347j - (AL021929) cobQ (Mycobacterium tuberculosis)		synthase	264602, 264769
471	17937351 (941, 942)	Novel Prolein sim. GBank gij114921 jspjP17447 jBETT_ECOLI - HIGH-AFFINITY CHOLINE TRANSPORT PROTEIN		transport	265019
472	R0047458 (943 944)				264596, 264685, 264557
473	20558793 (945, 946)	Novel Protein sim. GBank gil882343 (L10908) - Gcap1 gene product IMus musculus	6	UNCLASSIFIED	264369
474	80593365 (947, 948)			UNCLASSIFIED	22278997, 264692, 264288
475	82454665 (949, 950)			UNCLASSIFIED	264907, 264908, 264511, 265009, 264762, 264448, 284636, 264638
476	94143857 (851, 952)	Novel Protein sim. GBank gil5453656 ref NP_006329.1 pGAC1 - glioma amplified on chromosome 1 protein (leucine-rich)	Contains protein domain (PF00560) - glycoprotein Leudne Rich Repeat	glycoprolein	65274572, 60432049, 264259, 264508. 52644045, 55812038, 264758, 265011. 264288, 264686, 52644229, 65274791, 264638, 264566
477	79175833 (953, 954)			UNCLASSIFIED	264636
478	78633483 (955, 956)			UNCLASSIFIED	264690, 264693
479	80189746 (957, 958)			collagen	264886, 35695855, 265008, 264631, 264910. 264632, 264638, 265018, 264369, 264909
480	79390729 (959, 860)	Novel Protein sim. GBank gil1127551 (U18939) - orf2 [Battrachocottus baikalensis]		mapolymerase	264369
<u>\$</u>	79624578 (961, 962)			UNCLASSIFIED	264693
482	83050611 (963, 964)	Novet Protein sim. GBank gij4063042 (AF068065) - GP900; mucin-like giycoprotein [Cryptospondium parvum]		UNCLASSIFIED	264909, 264686, 264768, 264693, 55611576, 56182323, 18108385

1 1	20293306 (965, 966)		Contains protein domain (PF00534) - Glycosyt transferases group 1		264600
484	11618046 (867, 968)	Novel Protein sim. GBank gij3450883 (AF083334) - fibroin [Antheraea pernyi]	)		264594
	80191234 (969, 970)				264369, 21906765, 22279000, 22279002
486	80059042 (971, 972)	Novel Protein sim, GBank gil5042272 emb CAB44526.1  - (AL078618) nuoF, NADH dehydrogenase subunit  Streptomyces coelicolor		dehydrogenase	264604
	11813339 (973, 974)				264638
488	91222383 (975, 876)	Novel Protein sim. GBank gi 5724778 gb AAC53522.2  - Contains protein (AF012273) tho-type GTPase-activating protein thoGAPX-1 RhoGAP domain [Mus musculus]	Contains protein domain (PF00620) - RhoGAP domain		264686, 66714117, 264768, 18108385, 55811576, 265006, 265008, 265009, 265019, 22279002, 264259, 18108370, 264907, 24754, 56182323, 264288, 264693
489	10867710 (977, 978)	Novel Protein sim. GBank gij3882223]dbjjBAA34471.11- (AB018294) KIAA0751 protein [Homo sapiens]		kinase	264639
490	95361124 (979, 980)	Novel Protein sim. GBank gil82091lpirIIA25494 -		collagen	22278996, 29331822, 29331828, 264107,
		hydroxyproline-rich glycoprotein - tomato (fragment)		_	264909, 264110, 265009, 264592, 264593, 60433356, 264288, 264693, 263974, 263976, 20281071, 60432113
491	80498412 (981, 982)	Novel Protein sim. GBank gij2894206jemb CAA17072  - (AL021840) hypothetical protein Rv3258c [Mycobacterium tuberculosis]		UNCLASSIFIED	264769
482	87421264 (983, 984)				264600
493	11692942 (985, 986)			UNCLASSIFIED	264638
494	87726604 (987, 988)	Novel Protein sim. GBank giţ5262605 emb CAB45743.1 j -		UNCLASSIFIED	264489, 35696286, 60432289, 29331828.
		(AL080150) hypothetical protein [Homo sapiens]			35696052, 264509, 264905, 264906, 264907, 264908, 264909, 264909, 264510, 264511, 265009,
					264810, 3363/402, 264/62, 264/64, 264/66, 264/66, 264/69, 264/69, 264688, 21906765, 21906769
					35695917, 265020, 264693, 33657109,
					264629, 35696423, 35695855, 264634, 264638
485	80028599 (889, 990)	Novel Protein slm. GBank gij2791517 jemb j CAA16054 j - (AL021246) hypothetical protein Rv2477c [Mycobacterium tuberculosis]	Contains protein domain (PF00005) - transport ABC transporter	transport	264602, 264682, 264638
498	78985624 (991, 992)	Novel Protein sim. GBank gij230281jpdbj1R69j - 434 Repressor (Amino-Terminal Domain) (R1-69)	Contains protein domain (PF01381) - Helix-tum-helix		264601, 265021
497	78949661 (993, 994)	Novet Protein sim. GBank		oxidase	265006
		gil129736jspjP28225jPDXH_ECOLI - PYRIDOXAMINE 5:- PHOSPHATE OXIDASE (PNP/PMP OXIDASE)			

		2 [Rattus norvegicus]	Carboxylesterases		264509, 264905, 264906, 264907, 264908, 264509, 264510, 264501, 265009, 264910, 264511, 265009, 264910, 264591, 33657402, 264758, 265009, 264010, 264600, 264601, 264605, 264683, 264764, 264766, 264683, 264769, 21906767, 33657023, 284693, 264628, 264629, 35696423, 264630, 264632, 264634, 264635, 264563, 264565, 264566, 14108365, 264563, 264565, 264566,
8	20438222 (997, 998)	Novel Protein sim. GBank gij97480[piri]S19739 - integral membrane protein - Rhodobacter capsulatus		UNCLASSIFIED	264605
Ξ	11076810 (999, 1000)				264605
13	418034 (1001, 1002)	13418034 (1001, 1002) Novel Protein sim. GBank gi[5708250 emb CAB52363.1  - (AL109747) putative integral membrane protein [Streptomyces coelicolor A3(2)]		UNCLASSIFIED	264688
08	021176 (1003, 1004)	58678 emb CAB38132.1  - te isomerase (Streptomyces	Contains protein domain (PF00342) - Isomerase Phosphoglucose Isomerase	somerase	22278996, 265011, 264602, 264605, 264635
20	20264483 (1005, 1006)		J	UNCLASSIFIED	264564
10	10887321 (1007, 1008)			UNCLASSIFIED	264687
8	95003068 (1009, 1010)			UNCLASSIFIED	264369
<u> </u>	454292 (1011, 1012)	-M_TETPY - CALMODULIN	Contains protein domain (PF00036) - struct EF hand	struct	265010
2	451598 (1013, 1014)	20451598 (1013, 1014) Novel Protein sim. GBank gi[2501069]sp[Q46127 SYW_CLOLO - TRYPTOPHANYL- TRNA SYNTHETASE (TRYPTOPHANTRNA LIGASE) (TRPPRS)		UNCLASSIFIED	264604
2	184 j 424 (1015, 1016)	79841424 (1015, 1016) Novel Protein sim. GBank gil466068 sp P34618 YO82_CAEEL - HYPOTHETICAL 33.8 KD PROTEIN ZK1236.2 IN CHROMOSOME III		UNCLASSIFIED	264908
Ξ	11776386 (1017, 1018)	-			264638
83	83373465 (1019, 1020)			UNCLASSIFIED	264687, 264639
91	525578 (1021, 1022)	16525578 (1021, 1022)			265007
2	1399484 (1023, 1024 <u>)</u>	Novel Protein sim. GBank gi 2497419 sp P55635 Y4RB_RHISN - PUTATIVE INTEGRASE/RECOMBINASE Y4RB		UNCLASSIFIED	264565
82	1457404 (1025, 1026)		Contains protein domain (PF00134) - cyclin Cyclin	cyclin	264683, 264689, 35696423, 264639
32	<b>9813805 (1027, 1028)</b>	79813805 (1027, 1028) Novel Protein sim. GBank gil1184790 (U46068) - von Ebner minor salivary gland protein [Mus musculus]		UNCLASSIFIED	29331830, 264909
32	79462591 (1029, 1030)	_			22278999, 264690
8	9862020 (1031, 1032)	Novel Protein sim, GBank gi[2127400 pir  S65770 - mallooligosyltrehalose trehalohydrolase - Arthrobacter sp.		amylase	264910
1					

265018, 264605, 264764, 264766, 264687, 264891, 264691, 264565	264487	52644507, 22278997, 22278998, 60432049, 264259, 52645080, 29331824, 66714117, 60424269, 29331824, 66714117, 60424269, 29331826, 35696052, 264905, 29331830, 66712502, 264511, 265007, 264591, 60432229, 33657402, 60433438, 21906754, 33109954, 52644296, 87168474, 87168559, 265017, 265018, 264604, 265019, 265021, 60170815, 38557023, 224692, 252645129, 33657109, 27486262, 27486264, 3569563, 18108370, 264629, 52644332, 264639, 83373044, 18108385, 56526486, 60432113	264769	264605	264905, 264768	264629	ATPase_associated 284092, 264596, 265011	264907	264758	264769	56182575, 265017, 265018	265019	264687	264555, 264556, 264557, 264558, 18108385	264488, 264490, 264259, 264592, 264760, 265021, 264690, 263976, 264558
isomerase	UNCLASSIFIED	dna_ma_bind		transport	kinase	UNCLASSIFIED	ATPase_associated		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	hydrolase	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED
		Contains protein domain (PF01388) -   dna_rna_bind ARID DNA binding domain			Contains protein domain (PF00294) - kinase pfkB family carbohydrate kinase										Contains protein domain (PF00622) - UNCLASSIFIED SPRY domain
95292994 (1033, 1034) Novel Protein sim. GBank gil2983605 (AE000725) - ribose 5 protein sim. GBank gil2983605 (AE000725) - ribose 5		8) Novel Protein sim. GBank gij5689365 dbj BAA83073.1  - (AB024075) B120 [Homo saplens]	(0	11076821 (1041, 1042) Novel Protein sim. GBank gi 1169126 sp P46839 CTPA_MYCLE - CATION- TRANSPORTING P-TYPE ATPASE A	80435060 (1043, 1044) Novel Protein sim. GBank gij1172869jsp P44331 RBSK_HAEIN - RIBOKINASE	Novel Protein sim. GBank gi 2132243 pir  S61028 -     hypothetical protein YPL236c - yeast (Saccharomyces     cerevistae)	80261805 (1047, 1048) Novel Protein sim. GBank gil4033608 dbj BAA35136  -   (AB012308) B2HC (Anthocldaris crassispina)	(0)	36627630 (1051, 1052) Novel Protein sim. GBank gil4106610[emb]CAA21365j - (AL031866) ORF42, len=386 aa , similarity to an aminotransferase, in P95957 Sulfolobus solfataricus, (401 aa), 33.1% identity in 393 aa overlap, Fasta scores: opt:468, E(): 8.5e-24, in Q64602 R. norvegicus, (425 aa), 28.6% identi	(4)	(9)	17936810 (1057, 1058) Novel Protein sim. GBank gij731088 spjP24215 UXUA_ECOLI - MANNONATE DEHYDRATASE (D-MANNONATE HYDROLASE)		(2)	90933444 (1063, 1064) Novel Protein sim. GBank gij5262640 emb CAB45758.1  -  (AL080170) hypothetical protein  Homo sapiens
95292994 (1033, 1034)	8491831 (1035, 1036)	91677886 (1037, 1036	79869188 (1039, 1040)	11076821 (1041, 1042	80435060 (1043, 1044	18356013 (1045, 1046	80261805 (1047, 1048	79810046 (1049, 1050)	36827630 (1051, 1052	80504729 (1053, 1054)	85484134 (1055, 1056	17936810 (1057, 1058	10887336 (1059, 1060	80226576 (1061, 1062)	90933444 (1063, 1064
517			220		225	523	524	525	526		528		230	531	532

87761531 (1065, 1066) Novel Protein sim. GBank gi 4883838 gb AAD31593.1 AF11229 - (AF112299) Integral	Novel Protein sim. GBank   gl 4883638[gb]AAD31593.1 AF11229 - (AF112299	) integral			264907, 264909, 264768, 35695917, 264630, 264555
Inner nuclear membrane protein MAN1 (Homo sapiens) 82368284 (1067, 1068) Novel Protein sim. GBank gil2995352[emb]CAA04606.11-	inner nuclear membrane protein MAN1 (Homo sapiens)   Novel Protein sim. GBank gi 2995352 emb CAA04606.1   M01206) pan1 Petrantomyces confinenti	<u> </u> _		UNCLASSIFIED	264905, 265011, 264601, 264602, 264605, 264762, 264768, 265020, 264693, 264636
(7509, 1070) Novel racey pept per paraphany control (729128) similar to cAMP-dependant protein kinase; cDNA EST EMBL:100719 comes from this gene; cDNA EST yk46548.3 comes from this gene; cDNA EST yk46548.5	(X29128) Similar to CAMP-dependent protein kinase; cDN EST EMBL:100719 comes from this gene; cDNA EST yk465d8.3 comes from this gene; cDNA EST yk465d8.3 comes from this gene; cDNA EST yk465d8.3 comes from this gene; cDNA EST yk465d8.5 tomes from this gene; cDNA EST yk465d8.3 comes from this gene; cDNA EST yk492f4.3 comes from this generic cDNA EST yk492f4.3 comes from this		Contains protein domain (PF00069) - ATPase_associated 264906 Eukaryotic protein kinase domain	TPase_associated	:64 <u>906</u>
78907207 (1071, 1072) Novel Protein sim. GBank gilz495428jsplP55757fYOHL_SERMA - HYPOTHETICAL 10.1 KD PROTEIN IN BIOA 5'REGION	! I≪	-	N .	reductase	18108376, 264905, 264906, 264907, 264909
94147448 (1073, 1074)		H			265008, 264605, 65274791
87821963 (1075, 1076) Novel Protein sim. GBank gil134920 sp P21997 SSGP_VOLCA - SULFATED SURFACE GLYCOPROTEIN 185 (SSG 185)	) Novel Protein sim. GBank gil134920 sp P21997 SSGP_VOLCA - SULFATED SURFACE GLYCOPROTEIN 185 (SSG 185)	<u> </u>	Contains protein domain (PF00595) - collagen PDZ domain (Also known as DHR or GLGF).		29331822, 29331824, 29331825, 29331826, 29331827, 264908, 52644045, 33657402, 265017, 264762, 264683, 264288, 264685, 21906765, 35695763, 264558, 60170394, 264559, 22279002
28396269 (1077, 1078) Novel Protein sim. GBank gi 2498433 sp Q12341 HAT1_YEAST - HISTONE ACETYLTRANSFERASE				histone	264602, 265019
79637077 (1079, 1080)	-	L			264693
Novel Protein sim. GBank (AB018303) KIAA0760 pro	Novel Protein sim. GBank gij3882241(dbj BAA34480.1 - (AB018303) KIAA0760 protein [Homo sapiens]		Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type	transcriptfactor	18108394, 22278997, 22278998, 264259, 264112, 265009, 33657402, 55812038, 52648317, 265017, 21906765, 264693, 55811578, 264635, 56526486, 264566
95295836 (1083, 1084) Novel Protein sim. GBank gil5042272 emb CAB44526.1  - (AL078618) nuoF, NADH dehydrogenase subunit  Streptomyces coelicolor				dehydrogenase	264910, 265018, 264689, 264638, 264486
79796290 (1085, 1086)	(9)	-		UNCLASSIFIED	264602, 264908
20437191 (1087, 1088) Novel Protein sim. GBank gil2791398jemb CAA15994  - (AL021184) hypothetical protein Rv1464 [Mycobacterium tuberculosis]	Novel Protein sim. GBank gil2791398 emb CAA15994  - (AL021184) hypothetical protein Rv1464 [Mycobacterium tuberculosis]	<u> </u>		UNCLASSIFIED	264605
80434504 (1089, 1090)	+	$\vdash$			264768, 264634, 264907, 264592, 264909
80249016 (1091, 1092) Novel Protein sim. GBank gj 4887211jgb AAD32237.1jAF14744 - (AF147449) penicillin bindina protein 18 Pseudomonas aerudinosal		i.E			264600, 264602, 21906765
11077563 (1093, 1094) Novel Protein sim. GBank 11077563 (1093, 1094) Novel Protein sim. GBank RNA POLYMERASE BETA CHAIN (TRANSCRIPTASE BETA: CHAIN (RNA POLYMERASE BETA' SUBUNIT)	M) Novel Protein sim. GBank gi[1350855]sp P19176]RPOC_PSEPU - DNA-DIRECTED RNA POLYMERASE BETA' CHAIN (TRANSCRIPTASE BETA' CHAIN (RNA POLYMERASE BETA' SUBUNIT)			mapolymerase	264604
82114936 (1095, 1096) Novel Protein sim. GBank gil2330021 (AF018250) - kinesin- related protein; KRP; Costal2 [Drosophila melanogaster]	16) Novel Protein sim. GBank gij2330021 (AF019250) - kine related protein; KRP; Costal2 [Drosophila melanogaster]	Ė		UNCLASSIFIED	264488, 264505, 264910, 264760, 264693, 264639, 264563, 264564

UNCLASSIFIED 264488, 65274572, 18108398, 22278995, 22278997, 22278998, 22278996, 22278999, 22278999, 22278999, 22278999, 22278999, 264259, 29331824, 68714117, 29331825, 29331826, 264910, 264910, 264592, 23557402, 33109954, 265017, 265018, 265019, 18109351, 26448, 264688, 264688, 264688, 21906768, 21906768, 21906769, 264691, 33657023, 264692, 264693, 85274620, 52645129, 33557109, 27466261, 27466262, 204637, 2046	264688	UNCLASSIFIED 264908, 264909, 264768	UNCLASSIFIED 264689, 264639, 264563		264906	າດ 264762	ranscriptfactor 264508, 264605, 284559	dehydrogenase 264488	UNCLASSIFIED 264602	UNCLASSIFIED 264634
CANCE		ON O	CNC	08) - głycopi	kinase	58) - transpo	transcr	dehydr	ONCE OF THE PROPERTY OF THE PR	ONCE
				Contains protein domain (PF00008) - giycoprotein EGF-like domain		Contains protein domain (PF00358) - transport phosphoenolpyruvate-dependent sugar phosphotransferase system, EIIA 1				
95421904 (1097, 1098) Novel Protein sim. GBank gil4337460 gb AAD18133  - (AF056195) neuroblastoma-amplified protein [Homo saplens]		8U439990 (1101, 1102) Novel Protein sim. GBank gi 3122893 sp P94985 SYFB_MYCTU - PHENYLALANYL- TRNA SYNTHETASE BETA CHAIN (PHENYLALANINE- TRNA LIGASE BETA CHAIN) (PHERS)		80106002 (1105, 1106) Novel Protein sim. GBank gil552087 (M33753) - crumbs protein [Drosophila melanogaster]	/ ve 183 / v (1107, 1108) Novel Protein sim. GBank gi 5019771 gb AAD37857.1 AF13326 - (AF133263) histidine protein kinase-response regulator hybrid protein CvgSY [Pseudomonas syringae pv. syringae]	78995347 (1109, 1110) Novel Protein sim. GBank gil 31515jspjP02908jPTGA_SALTY - PTS SYSTEM, GLUCOSE-SPECIFIC IIA COMPONENT (EIIA-GLC) (GLUCOSE-PERMEASE IIA COMPONENT) (PHOSPHOTRANSFERASE ENZYME II, A COMPONENT)	20457127 (1111, 1112) Novel Protein sim. GBank gij3914014[splP96380]MFD_MYCTU - TRANSCRIPTION- REPAIR COUPLING FACTOR (TRCF)	19523405 (1113, 1114) Novel Protein sim. GBank gil5042273 emb CAB44527.1  -  AL078518) nuoE, NADH dehydrogenase subunit  Streptomyces coelicolor	20/24428 (1115, 1116) Novel Prolein sim. GBank gij11709331spP45331IMETE_HAEIN - 5- METHYLTETRAHYDROPTEROYLTIGLUTAMATE— HOMOCYSTEINE METHYLTRANSFERASE (METHIONINE SYNTHASE, VITAMIN-B12 INDEPENDENT ISOZYME) (COBALAMIN-INDEPENDENT METHIONINE SYNTHASE)	80084353 (1117, 1118) Novel Protein sim. GBank gil4980567[gb]AAD35173.1JAE00189 - (AE001694) fron(III)
95421904 (1097, 1098)	10886616 (1099, 1100)	80439990 (1101, 1102)	94672870 (1103, 1104)	80106002 (1105, 1106)	79618379 (1107, 1108)	78986347 (1109, 1110)	20457127 (†111, 1112)	19523405 (1113, 1114)	20/24429 (1115, 1116)	80084353 (1117, 1118)
	250		Š							228

			1 100000000		ADAMONDE NEADOR NEADON NEADOR
280	80066533 (1119, 1120)	80066533 (1119, 1120) Novel Protein sim. GBank Joil2402595lspiO531931Y4TR_RHISN - PROBABLE	ABC transporter		18108374
		DE ABC TRANSPOI			
		741K		INCI ASSIEIED	264600
100	20293187 (1121, 1122)				264680
562	(11698161 (1123, 1124)			טוגניטייטייטייטייטייטייטייטייטייטייטייטייטי	20400
263	79761420 (1125, 1126)	79761420 (1125, 1126) Novel Protein sim. GBank gil4104925 (AF042276) - poly(hydroxyalcanoate) granule associated protein GA2 [Pseudomonas putlda]		1	264910, 264691
564	56716390 (1127, 1128)	를 를		enase	264592
565	56465618 (1129, 1130)	56465618 (1129, 1130) Novel Protein sim. GBank gij349294(db) BAA32462  - (AB011532) MEGF6  Rattus norvegicus]	Contains protein domain (PF00008) - synthase EGF-like domain		265010
999	94323888 (1131, 1132)	94323888 (1131, 1132) Novel Protein sim. GBank gil4539568jemb CAB38487.1  -		helicase	264909, 264510, 265008, 284910, 264758.
		(AL035636) putative helicase [Streptomyces coelicolor]			264600, 284602, 284604, 284605, 284768, 284687, 264689, 35695917, 264693, 65274620, 264486
267	79560955 (1133, 1134)			UNCLASSIFIED	264681, 264691, 264593
568	94681793 (1135, 1136)	94681793 (1135, 1136) Novel Protein sim. GBank gij100506jpirj S17455 - Malate decarboxylating) (NADP+) dehydrogenase (oxaloacetate-decarboxylating) (NADP+) (FC 1 1 40) - Flaueria Innervia (frament)	Contains protein domain (PF00390) - dehydrogenase Malic enzyme	dehydrogenase	264689
	10077	Contract of the second which the second of t	Castains aratain domain (PEOD318) - Cibosomaintol	ribosomalorot	264565
699	38506897 (1137, 1j38)	39506897 (1137, 1138) Novel Protein sim. Gbank gil3915843 sp 031212 RS2_STRCO - 30S RIBOSOMAL PROTEIN S2	Ribosomal protein S2		
570	78375927 (1139, 1140)			UNCLASSIFIED	18108376, 18108387, 264565
1	70702064 (4444 4442)	70703084 (1444 4443) Main Destain eim CBank		Iransport	264907, 264909
5	721, 1141, 1147, 1147,	NOVERTOGEN SUN. GEGIN gil15122 sp 221627 BRAD_PSEAE - HIGH-AFFINITY BRANCHED-CHAIN AMINO ACID TRANSPORT PROTEIN BRAD			
572	36996838 (1143, 1144)			UNCLASSIFIED	264762
573	20715521 (1145, 1146)	20715521 (1145, 1146) Novel Protein sim. GBank gij4539223jembjCAB39881.1j - (AL049497) putative integral membrane protein		UNCLASSIFIED	265007, 264601
574	13521592 (1147 1148)	Suepionyes coencolor			264638
5	13076416 (1149 1150)	13076416 (1149 1150) Novel Protein sim. GBank		polymerase	264687
}		git 18794 sp  10443 DP3A_ECOLI - DNA POLYMERASE III. ALPHA CHAIN			
576		20482246 (1151, 1152) Novel Protein sim. GBank gij5457625jemb CAB49116.1 - (A_1248283) PAB2227 [Pyrococcus abyssi]			264605
577	66727102 (1153, 1154)	66727102 (1153, 1154) Novel Protein sim. GBank gi 5042274 emb CAB44528.1  -	Contains protein domain (PF00346) - dehydrogenase	- dehydrogenase	35696052, 264636
		(AL078618) nuoD, NADH dehydrogenase subunit [Strentomyces coelicator]	Respiratory-chain NADH dehydrogenase, 49 Kd subunit		•
578	11804477 (1155, 1156)	₹-			264638
579	Т	-		transport	264682, 264556
		gi 1723081 sp Q11046 Y089_MYCTU - HYPOTHETICAL ABC TRANSPORTER ATP-BINDING PROTEIN CY50.09			

					0337 00 4400000 000000000
580	80059417 (1159, 1160)				222 (899, 33030032, 204333, 204330, 284558
581	79230833 (1161, 1162)			UNCLASSIFIED	265008, 264564
582	80049817 (1163, 1164)	80049817 (1183, 1184) Novel Protein sim. GBank gij3243131 (AF045777) - titin IDrosophila melanogasteri	Contains protein domain (PF00047) - struct	struct	265021, 264555, 264557
583	79321392 (1165, 1166)	79321392 (1165, 1166) Novel Protein sim. GBank gil2501162 sp P77726 YAJR_ECOLI - HYPOTHETICAL 49,0 KD PROTEIN IN ABPA-CYOE INTERGENIC REGION		Iransport	264594
584	79845024 (1167, 1168)			UNCLASSIFIED	264488, 264908, 264766, 264687, 35696423
585	79581454 (1169, 1170)	79581454 (1169, 1170) Novel Protein sim. GBank gij3882221 dbjjBAA34470.1 -  (AB018293) KIAA0750 protein [Homo sapiens]		UNCLASSIFIED	265018, 264684, 21906769
588	38277486 (1171, 1172)			UNCLASSIFIED	264908, 265007
287	80497359 (1173, 1174)	80497359 (1173, 1174) Novel Protein sim. GBank gil4467250 emblCAB375751 -		hydrolase	264600, 264602, 264605, 264769, 264690.
		(AL035569) probable Glu-IRNA Gin amidotransferase subunit [Streptomyces coelicolor]			264557
288	79557239 (1175, 1176)	79557239 (1175, 1176) Novel Protein sim. GBank gil5689519 dbj BAA83043.1  -   (AB028014) KIAA1091 protein (Homo sapiens)		UNCLASSIFIED	265020, 264692
589	79805828 (1177, 1178)			UNCLASSIFIED	22278996, 264907, 264909, 264510, 265009,
					265010, 264687, 264769, 35695917, 18108376, 264634, 264638, 264638
280	79815629 (1179, 1180)			UNCLASSIFIED	264906, 264909
591	10313540 (1181, 1182)	10313540 (1181, 1182) Novel Protein sim. GBank gij2143293 embjCAB09390  - IZ355972) moB iMvcobacterium tuberculosis		mapolymerase	264691
265	13889767 (1183, 1184)	-		MHC	263972
593	82348699 (1185 1188)	Novel Protein sim GBank git4511983tablAAD21543.11 -		dehydrogenase	264511, 264762, 264769, 264486
}		(AF088896) electrotransfer ubiquinone oxidoreductase (IZymomonas mobilis)			
<u>8</u>	20212392 (1187, 1188)	20212392 (1187, 1188) Novel Protein sim. GBank gij1272368 (U51896) - LígE Nihdo parahaemolviicus		UNCLASSIFIED	264605
585	10064064 (1189, 1190)	1			264769
	,	gil131490 sp P20966 PTFB_ECOLI - PTS SYSTEM, FRUCTOSE-SPECIFIC IIBC COMPONENT (EIIBC-FRU)			
		(FRUCTOSE-PERMEASE IIBC COMPONENT) (PHOSPHOTRANSFERASE ENZYME II, BC COMPONENT) (FILFRII)			
596	13085170 (1191, 1192)	_		UNCLASSIFIED	264636
597	80259003 (1193, 1194)			UNCLASSIFIED	264592
298	94140216 (1195, 1196)	0		UNCLASSIFIED	264758, 55810764, 264555, 264558, 264637, 83373044
665	20385137 (1187, 1198	20385137 (1187, 1188) Novel Protein sim. GBank gif125328jsp P04951 KDSB_ECOLI - 3-DEOXY-MANNO- OCTULOSONATE CYTIDYLYLTRANSFERASE (CMP-KDO SYNTHETASE) (CMP-2-KETO-3-DEOXYOCTULOSONIC		UNCLASSIFIED	264603
Ş	10357663 (1100 1200)	ACID SYNTHETASE) (CKS)			264906
8	79610404 (1201, 1202	79610404 (1201, 1202) Novel Protein sim. GBank gil2127414lpirilS60064 -		UNCLASSIFIED	264510

	20632843 (1235, 1236)	20532843 (1235, 1235) Novel Protein sim. GBank gi[5459388]emb CAB50746.1] - (AL096839) putative aminotransferase [Streptomyces coelicolor]		isomerase	Z646U3
619	91227224 (1237, 1238)				56994075, 29331826, 33656970, 265008, 33657402, 33109954, 87188559, 264448, 18108374, 83373044
620	81183143 (1239, 1240)	81183143 (1239, 1240) Novel Prolein sim. GBank gil464335[sp]Q05922[DUS2. MOUSE - DUAL SPECIFICITY PROTEIN PHOSPHATASE 2 (DUAL SPECIFICITY PROTEIN PHOSPHATASE PAC-1)	,		29146498, 264758, 264369, 29148627
621	80239251 (1241, 1242)				264556, 264558, 264639
622	20456427 (1243, 1244	20456427 (1243, 1244) Novel Protein sim. GBank gil2633557 emb CAB13060  - (Z99110) yjdF (Bacillus subtilis)		SSIFIED	264605
623	10131798 (1245, 1246	10131798 (1245, 1246) Novel Protein sim. GBank gil1857710jgbjAAB482j -   C   (U87224) contactin associated protein [Rattus novegicus]	Contains protein domain (PF00054) - laminin Laminin G domain	laminin	264906
624	19534127 (1247, 1248	19534127 (1247, 1248) Novel Protein sim. GBank gil1705703 sp P52225 CCMF_PSEFL - CYTOCHROME C- TYPE BIOGENESIS PROTEIN CYCK	,	cylochrome	264596
. 625	13084619 (1249, 1250	13084619 (1249, 1250) Novel Protein sim. GBank gi[2894252[emb]CAA17114.1] - (AL021841) hypothetical protein Rv3342 [Mycobacterlum luberculosis]		UNCLASSIFIED	264688
929	88062603 (1251, 1252	n sim. GBank IP32323JAGA NT SUBUNIT		UNCLASSIFIED	29331822, 264905, 264908, 33657023, 33657109, 264558
627	80255457 (1253, 1254	80255457 (1253, 1254) Novel Protein sim. GBank gij3098418 (AF040944) - P140 [Mus musculus]		UNCLASSIFIED	18108394, 264112, 264593, 265022, 264635
628	80077096 (1255, 1256	m. GBank 50526 SSP1_SCHPO - DNINE-PROTEIN KINASE SSP1	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	264600
629	79851602 (1257, 1258	79851602 (1257, 1258) Novel Protein sim. GBank gij1143204 (U34305) - ORF2: Method: conceptual translation supplied by author. (Shigella sonnel)		isomerase	264906, 264907
630	39565156 (1259, 1260	39565156 (1259, 1260) Novel Protein sim. GBank gij3236368 (AF064748) - S3-12 [Mus musculus]		UNCLASSIFIED	264490
631	20598718 (1261, 1262	20598718 (1261, 1262) Novel Protein sim. GBank gil140687lsp P11666 YGGB_ECOLI - HYPOTHETICAL 30.9 KD PROTEIN IN SBM-FBA INTERGENIC REGION (ORF 4) (F286)			263978
632	27843890 (1263, 1264)			UNCLASSIFIED	264906, 264600, 264605, 264769, 264689, 264486
633	80477772 (1265, 1266)	10		UNCLASSIFIED	264769
ğ	17938806 (1267, 1268)	9)			265019
33	79574506 (1269, 1270)	0)		UNCLASSIFIED	264689
938	[79910981 (1271, 1272)	2)[		UNCLASSIFIED	264596, 264762, 264693

18 3	82455796 (1273, 1274) Novel Protein sim. GBank gi[2326739 emb CAB10953  -		nuclease	264906, 264907, 264510, 264511, 264601,
18268) recn (n	(298288) recn (Mycobacterium tuberculosis)			264762, 264766, 264687, 264769, 264689, 35695917, 264683, 264634, 264638, 264639, 264639, 264639, 264639, 264639, 264638, 264639, 264638, 264638, 264639, 264638, 264639, 264638, 264639, 264639, 264638, 264639, 264638, 2646888, 264688, 264688, 264688, 264688, 264688, 264688, 264688, 264
vel Protein 049645) p	14997457 (1275, 1276) Novel Protein sim. GBank gild678662 emb CAB41074.1  - (AL049645) putative large ATP-binding protein			264636
vel Protein	80204210 (1277, 1278) Novel Protein sim. GBank gil4589628 dbj BAA76836.1  -   Novel RAR02309 KIAA0992 protein Homo saninsi		struct	284112, 263974
vel Protein Skp1p, an	17929579 (1279, 1280) Novel Protein sim. GBank gill 432083 (U60981) - homolog Contains protein Contains by Conserved kinetochore protein in Skp1 family hydding yeaet (Arahidmests thaliana)	Contains protein domain (PF01466) - rnapolymerase Skp1 family	rnapolymerase	265009, 265010
			UNCLASSIFIED	264693
			UNCLASSIFIED	264565
			UNCLASSIFIED	264906, 264908, 264758, 264288, 264632, 264635, 264639, 264564
			UNCLASSIFIED	264684
			UNCLASSIFIED	264906, 264762, 264693, 264639, 264559
				264595, 264566
vel Prote 4507613	80257085 (1293, 1294) Novel Protein sim. GBank qil4507813IrefiNP 003738 1lpTNKS - TANKYRASE	Contains protein domain (PF00023) - transcriptfactor Ank repeat	transcriptfactor	264909, 264591
ovel Prote	80077428 (1295, 1296) Novel Protein sim. GBank gil1044963 bbs 169646 - protamine [Monodonta turbinata, gonads, Peptide, 106 aa]		UNCLASSIFIED	264600
			UNCLASSIFIED	263978
			UNCLASSIFIED	264686
Novel Prote gij1346916 ADENYLOS	11776932 (1301, 1302) Novel Protein sim. GBank gij1346916 sp P12283 pURA_ECOLI - ADENYLOSUCCINATE SYNTHETASE (IMP-ASPARTATE	4	·	264602, 264638
(100)			UNCLASSIFIED	264905, 264907, 264909, 263978, 264637
ovel Prot	82124947 (1305, 1306) Novel Protein sim. GBank		UNCLASSIFIED	22278996, 264510, 264511, 264512, 264593,
1722977 2.8 KD PF	gij1722977jsp Q10638 Y03C_MYCTU • HYPOTHETICAL 82.8 KD PROTEIN CY130.12C			21906754, 264603, 264760, 18108376, 264556
			UNCLASSIFIED	264906, 264595, 264632
ovel Prote 1130327]; LYCERO CYLTRAI CID ACY	79320692 (1309, 1310) Novel Protein sim. GBank gij130327jspjP26647jPLSC_ECOLI - 1-ACYL-SN-GLYCEROL-3-PHOSPHATE ACYLTRANSFERASE (1-AGP ACYLTRANSFERASE) (1-AGPAT) (LYSOPHOSPHATIDIC ACID ACYLTRANSFERASE) (LPAAT)	Contains protein domain (PF01553) - transferase Acyltransferase	transferase .	264592
			UNCLASSIFIED	264602, 264605, 264766, 264691
			LINCI ASSIFIED	284557 284558

								-									_
22278996, 60432049, 29331822, 29331824, 29331828, 265007, 265009, 33657402,	33657084, 265017, 264448, 21906765. 21906766, 263967, 20281149, 18108370. 18108374, 264482	72278999, 60432049, 66714117, 29331827, 265007, 264766, 56181562, 18108359, 18108365, 18108370, 18108381	264757	264510, 265009, 264600, 264602, 264603, 264604, 264605, 32833986, 18108376, 264636, 18108387, 22279000	264605	264605	264769	264600	264907, 265007	284689, 264602, 264593	264905, 264906, 264908, 264600, 264601, 264603, 264605, 264760, 264689, 264636, 264638, 264639	264637	264905, 264906, 264908, 264601, 264762, 264766, 264689, 264638, 18108385, 264486	264690, 264692, 264693, 264636, 18108387	264910, 265017	264759	264629
UNCLASSIFIED		transcriptfactor	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	dehydrogenase	UNCLASSIFIED	isomerase	glycoprotein	UNCLASSIFIED	ribosomalprot	UNCLASSIFIED	synthase	- UNCLASSIFIED	transcriptfactor		UNCLASSIFIED
		Contains protein domain (PF00096) - Iranscriptfactor Zinc finger, C2H2 type		Contains protein domain (PF00874) - UNCLASSIFIED Transcriptional antiterminator bgIG family							Contains protein domain (PF00453) - inbosomatprot Ribosomal protein L20			Contains protein domain (PF00023) - UNCLASSIFIED Ank repeat			
87761915 (1315, 1316) Novel Protein sim. GBank gi[5689493 db] BAA83030.1	Carolidae o mortina	87718663 (1317, 1318) Novel Protein sim. GBank gil2137872[pir] 148724 - zinc (Inger protein PZF - mouse		80026023 (1321, 1322) Novel Protein sim. GBank gij134180 sp P15401 SACY_BACSU - LEVANSUCRASE AND SUCRASE SYNTHESIS OPERON	20463731 (1323, 1324) Note of the sim CBank glass (AF11618) Sec glass (AF11618) Sec browners (AF11618) Sec browners (AF11618) Sec browners (AF1618) Sec br	20628080 (1325, 1326) Novel Protein sim. GBank gil5689250[db][BAA82881.1} - [Abovel 2433] similar to ori5 (Comamonas lestosteroni)	80508512 (1327, 1328) Novel Protein sim. GBank gil1652848įdbijBAA17766j - IO90909i DNA oholowase (Svnechocystis sp.)		78603142 (1331, 1332) Novel Protein sim. GBank gij3261829jembjCAB10927j - (298260) hypothetical protein Rv1230c (Mycobacterium Inherprofesis	94631802 (1333, 1334) Novel Protein sim. GBank gil5688851 dbj BAA82702.1 -	82051891 (1335, 1336) Novel Protein sim. GBank gi[3581853 emb CAA20809  - (AL031541) 50S ribosomal protein L20 [Streptomyces	_	80238549 (1339, 1340) Novel Protein sim. GBank gi 2582531 (AF026444) - 2-		79834371 (1343, 1344) Novel Protein sim. GBank gi[2114430 (U92703) - Olf-1/EBF- like-3 transcription (artor Mins minscribts)	82285798 (1345, 1346) Novel Protein slm. GBank. g  4589285  gb 4AD26430.1 AF13515 - (AF135154) ferric	alcargin studiopriore receptor (borrossia)
37761915 (1315, 1316)		87718663 (1317, 1318)	81897922 (1319 1320)		20463731 (1323, 1324)	20628080 (1325, 1326)	80508512 (1327, 1328)	80079053 (1329, 1330)	78603142 (1331, 1332)	94631802 (1333, 1334)	82051891 (1335, 1336)	12967154 (1337 1338)	80238549 (1339, 1340)	78601368 (1341, 1342)	79834371 (1343, 1344)	82285798 (1345, 1346)	79199259 (1347, 1348)
858 8		629	T	861	662	663	664	665	999	667	899	688	070	67.1	672	673	674

264488, 22278999, 66714117, 264508, 264511, 265008, 60433438, 264600, 264601, 264602, 264603, 264604, 264605, 264762, 264687, 264769, 60431602, 18108374, 264636, 264638	265010		264758, 264582, 264557	22278996, 264693	264600, 264602	264102, 264907, 264908, 265006, 264693, 263972, 83373044, 264568	264595	264634	264605	264909, 263967, 263981	264631	264693, 264635	264907, 264638	264906, 264907, 265007, 265009, 60433438.	21906/54, 264/50, 18106358, 21906/50, 21906/59, 265021, 18108361, 263974, 18108379, 264557, 18108385, 22279002	264510, 264511, 264764, 264769	264757	55812038, 55811957, 265018, 55811150, 18108351, 264908, 60431528, 264594	264635	22278996, 264510, 264602, 264603, 264762, 264687, 264769, 264693
JNCLASSIFIED		ATPase_associated	laminin	UNCLASSIFIED		laminin	UNCLASSIFIED	phosphalase					peptidase	UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	
Contains protein domain (PF01820) - UNCLASSIFIED D-ata D-ata ligase			Contains protein domain (PF00053) - laminin Laminin EGF-like (Domains III and V)			Contains protein domain (PF00053) - Iaminin Laminin EGF-like (Domains III and V)		Contains protein domain (PF00782) - phosphatase Dual specificity phosphalase,						Contains protein domain (PF00076) - UNCLASSIFIED	RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)			Contains protein domain (PF00097) - UNCLASSIFIED Zinc finger, C3HC4 type (RING finger)		*
1/AE00170 - (AE001708) D. Thermotoga maritima]	Novel Protein sim. GBank gij1723566isp Q10479 YDF7_SCHPO - PUTATIVE GLUCOSYLTRANSFERASE C17C9.07	587208 (U03976) - dynein teustes gratilla]	03365A -	78868855 (1357, 1358) Novel Protein sim. GBank gil3928723 emb CAA22219  - (AL034355) putative ABC transporter (Streptomyces coelicolor)		).1pLAMC - laminin, gamma 3		Novel Protein sim. GBank gil4758208jreftNP_004081.1pDUSP - dual specificity	phospinatase 3 (vacuna vius prospinatase virtiveaece) 20465367 (1367, 1368) (Novel Protein sim GBank 1915420319/emb(AB6679.1) -	Colored to the colore			Novel Protein sim. GBank gil2688962 (AF027768) - LspA (Serratia marcescens)	88064256 (1377, 1378) Novel Protein sim. GBank gij3046931 (AF049330) - PPAR	gamma coactivator [Mus musculus]	80389750 (1379, 1380) Novel Protein sim. GBank gi[2498941[sp]Q15428 SP62_HUMAN - SPLICEOSOME	ASSOCIATED THOTEIN OF (SAF OF) (STAND)	83608936 (1383, 1384) Novel Protein sim. GBank gi[5420387 emb CAB46679.1  - (AJ243459) proteophosphogiycan [Leishmania major]	78586116 (1385, 1386) Novel Protein sim. GBank gil854065 emb CAA58337  - I/X834131 U88 IHuman hemesvirus 61	82455983 (1387, 1388) Novel Protein sim. GBank gi 287327 sp Q01033 VG48_HSVSA - HYPOTHETICAL GENE 48 PROTEIN
87895870 (1349, 1350) Novel Protein sim. GBank gil4980755[gb AAD35347. alanineD-alanine ligase i	78899607 (1351, 1352) Novel Protein sim. GBank gij1723566 sp Q10479 YD GLUCOSYLTRANSFERA:	21644312 (1353, 1354) No	84225200 (1355, 1356) Nr lar	78868855 (1357, 1358) No. (A	20726424 (1359, 1360)		11302478 (1383 1384)		20465367 (1367, 1368) N		79208608 (1371 1372)	RD085629 (1373 1374)	79853412 (1375, 1376) N	88064256 (1377, 1378) N	<u>.</u>	80389750 (1379, 1380) h	19854 1987 /1381 13821	83608936 (1383, 1384)	79586116 (1385, 1386) I	82455983 (1387, 1388) 1
875 8	676 7	677	678	629	Г	681	T	683	684	200	88	Sa7	888	689		069	100	692	693	694

FR182575 264509 264905 264907	29331830, 264908, 264909, 264511, 265007, 264910, 264758, 264764, 264288, 65274791	264905, 264595	264682	264688, 35695917	264682	264693	264488, 264510, 264511, 264602, 264605, 264689	264909	264604	264288, 56181562, 33657109, 264629, 55811576	265006	264600	264603	264601, 264692	18108398, 264637, 264908, 264909	264369	265019	264591	264906, 264907	264508, 264555
INCI ACCIETED		transport	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	1	UNCLASSIFIED	UNCLASSIFIED	oxidase	UNCLASSIFIED	UNCLASSIFIED	,	transport		struct			- apolipoprolein	hydrolase	
Contract actions (DEOOOD)	4466339 emp CA636039.1 -	Contains protein domain (PF00005) - transport ABC transporter	Contains protein domain (PF01836) - UNCLASSIFIED Transposase							Contains protein domain (PF01344) - UNCLASSIFIED Kelch motif		,						Contains protein domain (PF00058)  Low-density lipoprotein receptor repeat dass B		
10 030000000000000000000000000000000000	94147849 (1389, 1390) Novel Protein sim. Gbank gil4468339 emplCA636039.1  -    (AJ010901) MUC4 [Homo sapiens]	79830982 (1391, 1392) Novel Protein sim. GBank gil2649950 (AE001058) - glutamine ABC transporter, ATP-binding protein (glnQ) [Archaeoolobus fulcidus]	11767889 (1393, 1394) Novel Protein sim. GBank gi 1731343 sp Q10694 yY25_MYCTU - HYPOTHETICAL 24 4 KD PROTEIN CY49.25				80230242 (1401, 1402) Novel Protein sim. GBank gil1001236 db  BAA10477  - (1084003) hypothetical protein (Synechocystis sp.)	+-	20446820 (1405, 1406) Novel Protein sim. GBank gi[2498935[sp]Q46338[SOXG_CORSP - SARCOSINE OXIDASE GAMMA SUBUNIT	94312224 (1407, 1408) Novel Protein sim. GBank gij3150513 (AF067219) - contains similarity to the kelch/MIPP family [Caenorhabditis	Novel Protein sim. GBank gil421091 pir  S30730 - hypothetical protein o206 - Escherichia coll	20288062 (1411, 1412) Novel Protein sim. GBank gi 3024872 sp Q55780 Y074_SYNY3 - HYPOTHETICAL 52 8 KD PROTEIN SI R0074	20838055 (1413, 1414) Novel Protein sim. GBank gij3420608jgbjAAC31907.1] - (AF075709) ABC transporter ATP-binding subunit   Pre-endomonas nutrida)	_	88001439 (1417, 1418) Novel Protein sim. GBank gij3649741jembjCAA03985j - (AJ000281) mucin [Homo saplens]	11356683 (1419, 1420) Novel Protein sim. GBank gij3080425jembjCAA18744.1j - (AL022604) puteiive protein fArabidoosis thalianal		80258164 (1423, 1424) Novel Protein sim. GBank gil4758686jrefjNP_002323.1jpLRP1 - low density lipoprotein Low-density lipoprotein receptor related protein 1 (alpha-2-macroglobulin receptor)	79263126 (1425, 1426) Novel Protein sim. GBank gij1703266 sp Q11056 AMI2_MYCTU - PUTATIVE	27847651 (1427, 1428) Novel Protein sim. GBank gi[4502351 ref NP_001692.1 pBAAT - bile acid Coenzyme A: amino acid N-acyltransferase; glycine N-
	94147849 (1389, 1390)	79830982 (1391, 1392)	11767889 (1393, 1394)	66695862 (1395, 1396)	79582558 (1397 1398)	79639098 (1399, 1400)	80230242 (1401, 1402)	78814789 (1403, 1404)	20446820 (1405, 1406	94312224 (1407, 1408	17932141 (1409, 1410)	20288062 (1411, 1412	20638065 (1413, 1414	20708292 (1415 1416)	88001439 (1417, 1418	11356683 (1419, 1420	17931418 (1421 1422)	80258164 (1423, 1424	79263126 (1425, 142	27847651 (1427, 142
П	589	969	269	869	Т	Т	701	702	703	704	705	706	707	708	60 `	5	1	712	713	714

264907	264692	264636	18108392, 22278994, 22278998, 265008, 265018, 264681, 18108354, 264684, 264685, 264686, 264687, 264689, 21906769, 18108361, 264691, 264692, 55810764, 264635, 18108382, 83373044, 18108388	265011	264908	264629	264910	264691	264909	264905, 264907	264636	264634	264567	264490	264564	264605	264591, 264594, 264595	264604	264604
UNCLASSIFIED		dehydrogenase		UNCLASSIFIED		UNCLASSIFIED		UNCLASSIFIED	protease	UNCLASSIFIED	UNCLASSIFIED			UNCLASSIFIED	isomerase	UNCLASSIFIED	struct	UNCLASSIFIED	ubiquitin
			Contains protein domain (PF00093) - von Willebrand factor type C domain						Contains protein domain (PF00353) - protease Hemotysin-type calcium-binding proteins										
79639423 (1429, 1430) Novel Protein sim. GBank gil1789035 (AE000352) - orf.		79491842 (1433, 1434) Novel Protein sim. GBank 79491842 (1433, 1434) Novel Protein sim. GBank 912494074[sp]P55653[GABD_RHISN - PROBABLE SUCCINATE-SEMIALDEHYDE DEHYDROGENASE (NADPA) (SSOH)	94319658 (1435, 1436) Novel Protein sim. GBank gij3873679[emb]CAA94886] - (Z71178) similar to pro-collagen domains; cDNA EST EMBL:D27978 comes from this gene; cDNA EST EMBL:D27977 comes from this gene; cDNA EST EMBL:D34199 comes from this g	17679564 (1437, 1438) Novel Protein sim. GBank gij2104302 emb CAB08631 - (295387) hypothetical protein Rv2611c  Mycobacterium (uberculosis)		15020180 (1441, 1442) Novel Protein sim. GBank gij123530jspjP04929jHRPX_PLALO - HISTIDINE-RICH GI YCOPROTEIN PRECURSOR	Novel Protein sim. GBank gl/498253 (U02372) - integrase	19755599 (1445, 1446) Novel Protein sim. GBank gi[2253054 emb]CAB10705  - (297559) Hipothetical protein Rv2114 [Mycobacterium theorytals]	10126494 (1447, 1448) Novel Protein sim. GBank gil4063015 (AF083061) - protease PrtA [Pseudomonas fluorescens]				20268471 (1455, 1456) Novel Protein sim. GBank gij2633910 emb CAB13411  - (299112) similar to hypothetical proteins [Bacillus subtilis]		19900373 (1459, 1460) Novel Protein sim. GBank gilz494660 sp Q45291 GALE_BRELA - UDP-GLUCOSE 4- EPINERASE (GALACTOWALDENASE) (UDP-	80058750 (1461, 1462) Novel Protein Sim. GBank gi[1146192 (L47838) - putative	80258175 (1463, 1464) Novel Protein sim. GBank gij1168396jspjP46681JAIP2_YEAST - ACTIN INTERACTING PROTEIN 2	_	20435987 (1467, 1468) Novel Protein sim. GBank gij3184080jembjCAA19336j - (AL023781) hypothetical protein [Schizosaccharomyces pombe]
9639423 (1429, 1430) N	0660000 (4404 4400)	9491842 (1433, 1434) N 9491842 (1433, 1434) N 9	4319658 (1435, 1436) N (c) E E	17679564 (1437, 1438) h ((	79841684 (1439, 1440)	15020180 (1441, 1442) h	9862603 (1443, 1444)	19755599 (1445, 1446)	10126494 (1447, 1448)	70878879 (1449 1450)	13086282 (1451, 1452)	13522872 (1453, 1454)	20268471 (1455, 1456)	11293753 (1457, 1458)	+	80058750 (1461, 1462)	80258175 (1463, 1464)	20446839 (1465, 1466)	20435987 (1467, 1468)
715 79	Т	717	718	719	720 7		722	723	724	725	Т	Г	1	Т	730	731	732	Т	75

1054504	704304	264636	265020	264910	254007 264764 264634 264637	264507, 264704, 204504, 204508, 264508, 264508,	18108362	10.00 10.00	264488, 264259, 264508, 264905, 264906, 264907, 264907, 264909, 264510, 264511, 264512, 265008, 265009, 264910, 264591, 264592, 264596, 265011, 18108351, 264769, 264766, 264769, 264769, 264693, 18108370, 264629, 18108372, 264630, 264631, 264639, 18108372, 264630, 264667	18108397, 264511, 264690, 264628, 264638, 264638, 264639, 264768	264906	264600	66714117, 264909, 264509, 264900, 264907, 264908, 264909, 264511, 264910, 265011, 264081, 264781, 264781, 264789, 264789, 264789, 264691, 264693, 264693, 264634, 264635, 264639, 26183323, 83373044	264690
	-	iransport	UNCLASSIFIED	o o o pino		INCI ASSIBILED		UNCLASSIFIED	UNCLASSIFIED		glycoprolein	synthase	- oxidase	
		Contains protein domain (PF00528) - transport Binding-protein-dependent transport systems inner membrane component	Contains protein domain (PF00615) - UNCLASSIFIED Regulator of G protein signaling domain			10070101	Contains protein domain (Pro 1865) - Orocassii isaa Domain of unknown function	,	Contains protein domain (PF00145) - UNCLASSIFIED C-5 cytosine-specific DNA methylase				Contains protein domain (PF00018) - oxidase SH3 domain	
	11607959 (1469, 1470) Novel Protein sim. GBank gil401582 sp P27432 YICE_ECOLI - HYPOTHETICAL 48.9	ECOLI - PUTRESCINE RMEASE PROTEIN POTH			79833670 (1477, 1478) Novel Protein sim. GBank gi[2506867]sp[P33225 TORA_ECOLI - TRIMETHYLAMINE- n-OXIDE REDUCTASE PRECURSOR (TMAO REDUCTASE) (TRIMETHYLAMINE OXIDASE)			82393795 (1483, 1484) Novel Protein sim. GBank gij3877494[emb]CAA88472.1] - (Z48583) ATP binding protein with similarity to the CDC48/PAS1/SEC18 family; cDNA EST EMBL:D65037 comes from this gene; cDNA EST EMBL:D68340 comes from this gene; cDNA EST EMBL:D68340 comes	gene; cDNA EST EMBL:D6845 82300051 (1485, 1486) Novel Protein sim. GBank gij127420jspjP19888jMTBA_BACAR - MODIFICATION METHYLASE BANI (CYTOSINE-SPECIFIC METHYLTRANSFERASE BANI) (M.BANI)	0	Novel Protein sim. GBank gi 78921 pir  S04846 - UDP-N-acetylmuramoylalanyl-D-glutamyl-2, 6-diaminopimelateD-alanine ligase (EC 6.3.2.15) precursor -	1073229 (1491, 1492) November 1 (1892)	94322044 (1493, 1494) Novel Protein sim. GBank gi[288741 db] BAA24848 - (AB007878) KIAA0418 [Homo sapiens]	16
	1607959 (1469, 1470) N	N (1471, 1472) N (1471, 1472) N (9)	78945340 (1473, 1474)	17895353 (1475, 1476)	<del></del>	19881557 (1479, 1480)	79827273 (1481, 1482)	52393795 (1483, 1484)	82300051 (1485, 1486)	80230421 (1487, 1488)	9841963 (1489, 1490)	11073229 (1491, 1492)	94322044 (1493, 1494)	30.1.
	735 11	736 10	737	807		97	$\top$	742	743	744	745	746	747	

Г	20469119 (1497, 1498)	498)			UNCLASSIFIED	264604
750	20296427 (1499, 15	2003 2003	20296427 (1499, 1500) Novel Protein sim. GBank		ł	264600
		<u> </u>	giji169727jspjP44946jFPG HAEIN - FORMAMIDOPYRIMIDINE-DNA GLYCOSYLASE (FAPY- DNA GLYCOSYLASE)			
751	21636169 (1501, 15	9 (305) N	21636169 (1501, 1502) Novel Protein sim. GBank gi 5360068 gb AAD42851.1 AF15968 - (AF159689) serineAhreonine kinase PKN3 [Myxococcus xanthus]	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	264605, 264559
752	82450366 (1503, 15	2 5 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	82450366 (1503, 1504) Novel Protein sim. GBank gil1168662[sp P44426 BIOA_HAEIN - ADENOSYLMETHIONINE-8-AMINO-7-OXONONANOATE AMINOTRANSFERASE (7,8-DIAMINO-PELARGONIC ACID AMINOTRANSFERASE) (DAPA AMINOTRANSFERASE)	Contains protein domain (PF00202) - UNCLASSIFIED Aminotransferases class-III pyridoxal phosphate	UNCLASSIFIED	264508, 35695855, 264638, 18108387
753	80508718 (1505, 15	2 8 8 3)	80508718 (1505, 1506) Novel Protein sim: GBank gi 2851530 sp P32399 YHGE_BACSU - HYPOTHETICAL 84.1 KD PROTEIN IN HEMY-GLTT INTERGENIC REGION (ORFB)		UNCLASSIFIED	264909, 264600, 264602, 264604, 264760, 264769, 264634
754	95083741 (1507, 1508)				UNCLASSIFIED	264508, 264906, 264907, 264908, 264909, 264759, 264602, 264604, 284769, 264628, 264629, 264639, 264634, 264635, 264637, 264637, 264637, 264637, 264638, 264637, 264638
755	80185449 (1509, 1510)	510			UNCLASSIFIED	264448, 264690
952	94631686 (1511, 15	1512) N	94631686 (1511, 1512) Novel Protein sim. GBank gij349276 emb CAA20420  - (AL031317) putative dehydrogenase (Streptomyces coelicotor)			264769, 264689, 264638, 264639
757	79468533 (1513, 1514)	1514)			UNCLASSIFIED	264682, 264685
758	78963176 (1515, 15	1516) N (4)	78983176 (1515, 1516) Novel Protein sim. GBank gil4580331 emb CAB40107.1  - (AJ001206) putative glycogen debranching enzyme [Streptomyces coelicolor]		amylase	265007, 18108387, 265007, 18108387
759	79475667 (1517, 19	1518) h	79475667 (1517, 1518) Novel Protein sim. GBank gi[2911858 (AF047659) - No   definition fine found [Caenorhabditis elegans]		UNCLASSIFIED	264684, 264686
760	87628888 (1519, 16	1520) h	87628888 (1519, 1520) Novel Protein sim. GBank gil3451312 emb CAA20449  - (AL031324) membrane atpase (Schizosaccharomyces pombe)	Contains protein domain (PF00122) - transport E1-E2 ATPase	transport	29331822, 264908, 52644045, 56182435, 60170831, 21906754, 265017, 265019, 264681, 264687, 264688, 21906766, 21906768, 265020, 265021, 265022, 264635, 22279000
781	79877966 (1521, 1522)	1522)			UNCLASSIFIED	264768
782	80023563 (1523, 16	1524)	80023563 (1523, 1524) Novel Protein sim. GBank gi[3327158 dbj BAA31647] -   (AB014572) KIAA0672 protein [Homo saplens]		UNCLASSIFIED	264907, 264593, 265020
763	20294813 (1525, 18		20294813 (1525, 1526) Novel Protein sim. GBank gil4981266igblAAD35822.1IAE00174 - (AE001744) lipopolysaccharide core biosynthesis protein KdtB [Thermotoga maritima]			264600
764	39515024 (1527, 1528)	-				264603

264905, 264906, 264594, 264686, 33657023			264592, 264595	55810764, 264559	J 264905, 264907, 264828, 264909, 265010.					D 264910				265019			264603	264907, 264600, 264601, 264602, 264603, 264604, 264605, 264486	264905, 264907, 265010, 264600, 264601, 18108362, 18108374, 264558		265020		35696052, 264604	264603	ED 264593
struct		UNCLASSIFIED	UNCLASSIFIED	esterase	UNCLASSIFIED		UNCLASSIFIE			UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED		protease	UNCLASSIFIED		synthase		UNCLASSIFIED		UNCLASSIFIED	struct	transport	UNCLASSIFIED
							Contains protein domain (PF00097) - UNCLASSIFIED		(inger)					Contains protein domain (PF01006) - Hepatitis C virus non-structural protein NS4a	Contains protein domain (PF01344) - protease Kelch motif								Contains protein domain (PF00047) - struct		
80025347 (1529, 1530) Novel Protein sim. GBank gij3845093 (AE001371) - erythrocyle membrane protein PfEMP3 (Plasmodium falcinanum)		10296742 (1533, 1534) Novel Protein sim. GBank gij541121 pir  540827 - hvpothetical protein o300 - Escherichia coll		80086554 (1537, 1538) Novel Protein sim. GBank gilz982501 jembiCAA061841 -	80417847 (1539 -1540) Novel Protein sim GBank di283437 loiri S27850 -	hypothetical protein - Trypanosoma cruzi (fragment)	95329509 (1541, 1542) Novel Protein sim. GBank	gil4769004[gbfAAD29715.1]AF14059 - (AF140598) ring-box	protein 1 [Homo sapiens]			79856129 (1547, 1548) Novel Protein sim. GBank gij5531324[embjCAB51045.1] - (AJ009579) putative alkane 1-monooxygenase (Pseudomanas fluorescens)	-		78950378 (1553, 1554) Novel Protein sim. GBank gij4505461 įretįNP_003624.1 įpNRPB - nuclear restricted protein. BTB domain-like (brain)			95288987 (1559, 1560) Novel Protein sim. GBank gij1144520 (U34956) - phosphoribosylformylglycinamidine synthase [Mycobacterium tuberculosis]		Novel Protein sim. GBank gild 155447 (AE001517) - proline/betaine transporter [Helicobacter pylori J99]	٠.		56073541 (1569, 1570) Novel Protein sim. GBank gij3451335 (AC005525) - F22162 1 Homo sapiensi	20438842 (1571, 1572) Novel Protein sim. GBank gijt38748jspjP10905jUGPA_ECOLI - SN-GLYCEROL-3- PHOSPHATE TRANSPORT SYSTEM PERMEASE PROTEIN LICPA	
80025347 (1529, 1530) [1	82417404 (1531 1532)	10296742 (1533, 1534)	79416080 (1535, 1536)	80086554 (1537, 1538)	90417847 (1539 -1540)		95329509 (1541, 1542)			78971362 (1543 1544)	78945383 (1545, 1546)	79856129 (1547, 1548)	20620141 (1549, 1550)	78942693 (1551, 1552)	79960378 (1553, 1554)	20691310 (1555, 1556)	80054024 (1557, 1558)	95288987 (1559, 1560)	80250049 (1561, 1562)	8758529 (1563, 1564)	16410791 (1565, 1566)	80051197 (1567, 1568)	56073541 (1569, 1570)	20438842 (1571, 1572)	80258364 (1573, 1574)
765	T	767	Т	769	27.0		771			772	Т		775	776	111	778	179	780	781	782	783	784	785	786	787

264909, 264602, 264603, 264769, 264638	265007	72278999, 29331824, 264828, 87168559, 265018, 21906765, 21906767, 21906768, 21906769, 265020, 264692, 22279000, 264563	264601, 264605, 264636	265021, 264631, 264635, 264536	264693	264603, 264604, 264910, 264762, 264900. 264639, 264909, 264757	2694488, 35696052, 264905, 264907, 255010. 35696423, 264636	60432229, 55811150, 264530, 254537, 264565	22278998, 264093, 264094, 66714117, 21906767, 21906769, 265020, 265022	264687, 264768, 264693	264909, 264910, 264639, 204630	264488	264508, 264511, 265006, 265009, 264769, 264567, 264486	264905, 264593	264605	264510	264508	264769	264764
synthase	UNCLASSIFIED	glycoprotein		struct	ротворох	UNCLASSIFIED	carboxylase		struct	transcriptfactor	hydrolase	UNCLASSIFIED	transport		struct	kinase		- transferase	
		Contains protein domain (PF00047) - glycoprotein Immunoglobulin domain			Contains protein domain (PF00046) - homeobox Homeobox domain													Contains protein domain (PF00185) - transferase Asparlate/ornithine carbamovitransferase	
80507844 (1575, 1576) Novel Protein sim. GBank gil2746079 (AF015310) - BTH1	17294715 (1577, 1578) Novel Protein sim. GBank gil2351849 (U93357) - 40 kDa	R6284408 (1579, 1580) Novel Protein sim. GBank gils706378[db][BAA83099.1] (AB026118) MALT1 [Homo saplens]	94651627 (1581, 1582) Novel Protein sim. GBank gil5689948jemb CAB51985.1 - (AL109863) putative isoleucyl-tRNA synthetase isoleucyl-tRNA synthetase	80058786 (1583, 1584) Novel Protein sim. GBank gij393194 (L02375) - S-antigen Protein sim. GBank gij393194 (L02375) - S-antigen falchoarumi	79638730 (1585, 1586) Novel Protein sim. GBank gil 1345408 db  BAA05046  -		80074988 (1589, 1590) Novel Protein sim GBank gill 87334 emb[CAB07082] - ISBO 1589, 1590) Novel Riveobacierum Weercaliosis!		87771781 (1593, 1594) Novel Protein sim. GBank gil2995447[emb[CAA71519] - 17910495; CDV-1R protein [Mus musculus]		79557816 (1597, 1598) Novel Protein sim. GBank gil4467250jemb CAB37575  - (AL035569) probable Glu-tRNA Gin amidotransferase	Suburin (Strepton) yes coencero)	80499399 (1601, 1602) Novel Protein sim. GBank gi[2791517]emb[CAA16054] - (AL021246) hypothetical protein Rv2477c [Mycobacterium tuberculosis]	79834598 (1603, 1604) Novel Protein sim. GBank gjl4887211[gb]AAD32237.1[AF14744 - (AF147449) penicillin bindina protein 18 [Pseudomonas aerudinosa]		10174239 (1607, 1608) Novel Protein sim. GBank gij1176152jspP44507jYHAD_HAEIN - HYPOTHETICAL PROTEIN HINGS		80484113 (1611, 1612) Novel Protein sim. GBank gil2764612 emb CA404683  - 80484113 (1611, 1612) Novel Protein sim. GBank gil2764612 emb CA404683  -	80381812 (1613, 1614) Novel Protein sim. GBank gi 2833311 sp 021828 YNFD_CAEEL- HYPOTHETICAL 18.9 KD PROTEIN R07E5,13 IN CHROMOSOME III
0507844 (1575, 1576) No	T294715 (1577, 1578) N	, 6284406 (1579, 1580) N	14651627 (1581, 1582) N (/	30058786 (1583, 1584) N	79638730 (1585, 1586) N	91839294 (1587, 1588) h	90074988 (1589, 1590) h	86669451 (1591, 1592)	87771781 (1593, 1594)	70865200 (1505 1506)		70070180 /1500 1500	80499399 (1601, 1602)	79834598 (1603, 1604)	20467520 (1605, 1606)	10174239 (1607, 1608)	7060003 /1600 16101	80484113 (1611, 1612)	80381812 (1613, 1614)
788 80	789 1	790 8	791	792 8	783	797	795	962	797	Т	789	Т	8 6 8	802	Т	804	4	908	807

1					
	35106817 (1615, 1616)	35106817 (1615, 1816) Novel Protein sim. GBank gij3913092 sp Q46170 ARCD_CLOPE - ARGININE/ORNITHINE ANTIPORTER		transport	264909, 264602, 21906764, 18108374
T	81454254 (1617, 1618)	81454254 (1617, 1618) Novel Protein sim. GBank gij3913016jspjP74309JALF1_SYNY3 - FRUCTOSE- BISPHOSPHATE ALDOLASE CLASS I (FBP ALDOLASE)	Contains protein domain (PF00274) - UNCLASSIFIED Fructose-bisphosphate aldotase dass-l	UNCLASSIFIED	264508, 264906, 264909, 265007, 264910, 264758, 264600, 264602, 264603, 264605, 264687, 264769, 264689, 264636, 264486
	80192761 (1619, 1620)	80192761 (1619, 1620) Novel Protein sim. GBank gil401472 sp P30863 YAFB_ECOLI - HYPOTHETICAL OXIDOREDUCTASE IN ASPU-MLTD INTERGENIC REGION	Contains protein domain (PF00248) - reductase Aldo/keto reductase family	reductase	284369
	80079280 (1621, 1622)			UNCLASSIFIED	264558
1	10297654 (1623, 1624)			UNCLASSIFIED	264692
1	79612280 (1625, 1626)			UNCLASSIFIED	264906
	80473427 (1627, 1628)	80473427 (1627, 1628) Novel Protein sim. GBank gi[146168 (J01617) - glutaminyl- tRNA synthetase [Escherichia coli]		synthase	264905, 264602, 264605, 264682, 264687, 264769, 264636
1 _	95419513 (1629, 1630)	95419513 (1629, 1630) Novel Protein sim. GBank gil4589652 db  BAA76848.1  -		UNCLASSIFIED	264488, 22278998, 22278999, 29331822,
_		(AB023221) KIAA1004 protein [Homo saplens]			29331824, 29331825, 29331827, 29331828, 29146499, 264905, 264908, 265007, 33657402, 60433356, 60433438, 264758, 265011, 265017, 265018, 265019, 264369, 264288, 264685, 21906765, 21906767, 265020, 265021, 264692, 65274620, 33657109, 264629, 18108376, 284635, 264638, 60170394, 56182323, 264584
	19881910 (1631, 1632)				264600
	95293316 (1633, 1634)	95293316 (1633, 1634) Novel Protein sim. GBank gi 1781144 emb CAB06254  - (Z83865) hypothetical protein Rv3069 [Mycobaclerium [tuberculosis]		UNCLASSIFIED	264595
	90938190 (1635, 1636)	90938190 (1635, 1636) Novel Protein sim. GBank gil1477468 (U35244) - vacuolar		-	65274572, 22278999, 60424269, 35696052,
1					55812038, 21906768, 55811957, 35695917, 33657023, 18108370, 18108374, 55810764, 35696423, 55811576, 264636
	80254977 (1637, 1638)	80254977 (1637, 1638) Novel Protein sim. GBank gi 1001352 dbj BAA10839  - (D64006) ABC transporter (Synechocystis sp.)		transport	264565
	80059688 (1639, 1640	Novel Protein sim. GBank gij586814 splP37484 YYB 74.3 KD PROTEIN IN RPL		UNCLASSIFIED	264600, 264602, 264604
	79762590 (1641, 1642)			UNCLASSIFIED	264910
	80215310 (1843, 1644)			UNCLASSIFIED	264510, 264594, 264637
	94992299 (1645, 1646	94992299 (1645, 1646) Novel Protein sim. GBank gij3878400 emb CAA95828  - (Z71284) predicted using Genefinder; Weak similarity to Mouse T-complex-associated-testes-expressed-1 protein (PIR Acc. No. A45841); cDNA EST EMBL:D32742 comes from this gene; cDNA EST EMBL:D33617 comes from this gene; cDNA EST.		struct	264509, 264687, 264691
	80411171 (1647, 1648	80411171 (1847, 1648) Novel Protein sim. GBank gi 1370076 emb CAA66887 -  (X98235) type I [Drosophila melanogaster]	Contains protein domain (PF01429) - Methyl-CpG binding domain		264910, 264763, 264769, 264693

	264605		52645080, 264508, 264905, 264509, 264906, 264907, 264908, 264909, 264512, 264910, 264591, 264591, 264560, 264768, 269659, 264631, 264628, 2564630, 264631, 264638, 264631, 264638, 264638, 264637, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264688, 264638, 264688, 264						264600, 264602, 264769, 264689, 264636	264768, 263994, 21906767, 264910, 264632, 264635, 264259, 264639, 264693, 83373044, 264758, 35696052, 22279002, 264908, 264906, 264448, 263972, 264908, 264909	35695917, 264557	D 264595		D 264906	pt 264762, 264556
UNCLASSIFIED			UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	kinase		transport	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	- interleukinrece
			•					·	Contains protein domain (PF00224) - kinase Pyruvate kinase	·					Contains protein domain (PF00097) - interleukinrecept Zinc finger, C3HC4 type (RING finger)
20638600 (1649, 1650) Novel Protein sim. GBank gij3025132[spl977391]YEAG_ECOLI - HYPOTHETICAL 74.5 KD PROTEIN IN GAPA-RND INTERGENIC REGION	11075047 (1651, 1652) Novel Protein sim. GBank gij3242281[emb CAA16669] - (AL021646) hypothetical protein Rv3202c [Mycobacterium tuberculosis]	80054207 (1653, 1654) Novel Protein sim. GBank gij3417424jemb CAA20312  - (AL031261) putative transport protein   Schizosaccharomyces pombel	95105322 (1655, 1656) Novel Protein sim. GBank gil4336992 gb AAD17897  - (AF101361) Abnormal X segregation [Drosophila melanogaster]		20395091 (1659, 1660) Novel Protein sim. GBank gij3820539 (AF080002) - UDP-N-acetylmuramyl tripeptide synthetase MurC [Heliobacillus mobilis]		19536322 (1663, 1664) Novel Protein sim. GBank gi 1970004 emb CAB06855  - (Z92539) hypothetical protein Rv1024 (Mycobacterium tuberculosis)	20726654 (1665, 1666) Novel Protein sim. GBank gil2500056lsplQ46267lPFLA_CLOPA - PYRUVATE FORMATE-LYASE ACTIVATING ENZYME	21428762 (1667, 1668) Novel Protein sim. GBank gil2497531 sp Q46078 KPYK_CORGL - PYRUVATE KINASE (PK)		66126552 (1671, 1672) Novel Protein sim. GBank gil699315 (U15184) - phosphate Itransport protein PSTC [Mycobacterium leprae]		79184203 (1675, 1676) Novel Protein sim. GBank gif728867\spIP40602\APG_ARATH - ANTER-SPECIFIC PROLINE-RICH PROTEIN APG PRECURSOR		80059851 (1679, 1680) Novel Protein slm. GBank gil4557753 ref NP_000372.1 pMID1 - midline 1 protein
20638600 (1649, 1650)	11075047 (1651, 1652)	80054207 (1653, 1654)	95106322 (1655, 1656)	81742215 (1657, 1658)	20396091 (1659, 1660)	87112435 (1661, 1662)	19536322 (1663, 1664)	20726654 (1665, 1666)	21428762 (1667, 1668)	94140482 (1669, 1670)	66126552 (1671, 1672)	79450450 (1673, 1674)	79184203 (1675, 1676)	79641125 (1677, 1678)	80059851 (1679, 1680)
825	826	827	828	829	830	831	832	833	834	835	836	837	838	839	940

2	80376318 (1681, 16	(289	80376318 (1681, 1682) Novel Protein sim. GBank gil139805(splP08045)XFIN XENLA - XFIN PROTEIN	Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type		264764
842	80078724 (1683, 16	( <del>8</del> 4	80078724 (1683, 1684) Novel Protein sim. GBank gil2114321 dbj BAA20037  - (D88733) membrane glycoprotein [Equine herpesvirus 1]	Contains protein domain (PF00569) - UNCLASSIFIED Zinc finger present in dystrophin, CBP/p300	UNCLASSIFIED	264905, 264908, 265008, 265009, 18108374, 56182323, 264558
843	87002847 (1685, 16	(989	87002847 (1685, 1686) Novel Protein sim. GBank gij3882225[dbjjBAA34522.1] - (AB018345) KIAA0802 protein [Homo saplens]	Contains protein domain (PF00170) - struct bZIP transcription factor	struct	264091, 29331825, 264906, 264768, 264563
844	17941439 (1687, 16	(88)	17941439 (1687, 1688) Novel Protein sim. GBank gij2224721 dbj BAA2084 - (AB002388) KIAA0390 [Homo saplens]	Contains protein domain (PF00096) - transcripifactor Zinc finger, C2H2 type		265011
845	18346844 (1689, 1690)	(069				264629
848	79863441 (1691, 16	(269	78863441 (1691, 1692) Novel Protein sim. GBank gi[625679 ptr  A36929 - virulence regulatory protein VsrB - Pseudomonas solanacearum		kinase	264907
847	78695348 (1693, 1694)	88				264909
848	78489365 (1695, 1696)	(969			UNCLASSIFIED	265020
849	79756367 (1697, 1698)	(869			UNCLASSIFIED	264566
850	79817849 (1699, 1700)	700)	Novel Protein sim. GBank gij3183245 sp P78061 YCJK_ECOLI - PUTATIVE GLUTAMINE SYNTHETASE (GLUTAMATE-AMMONIA LIGASE)	Contains protein domain (PF00120) - UNCLASSIFIED Glutamine synthetase	UNCLASSIFIED	264909
			gi 5454130 ref NP_006280.1 pTLN  - talin	MLWEQ domain		65274577, 50182576, X2278997, Z2278998, Z2278996, 22278999, 22278999, 222278999, 222278999, 222278999, 222278999, 222278999, 222278999, 222278999, 222278999, 222278999, 222278999, 222378297, 22228996, 29331824, 66714117, 265406, 29331824, 66714117, 265406, 29331826, 29331827, 2644045, 56182435, 265006, 265007, 265008, 265009, 264910, 60432229, 60431735, 60433356, 33657402, 60433738, 264599, 264760, 265011, 265018, 265019, 264760, 264761, 264762, 264619, 18108351, 264760, 264761, 264762, 26488, 264762, 26488, 264766, 264766, 264769, 264769, 29148629, 55811957, 35695917, 265020, 265021, 265022, 60170615, 52644150, 264691, 264692, 33657023, 264693, 264692, 33657023, 264693, 263666, 236526, 2486522, 2486526, 248626, 2486526, 248626, 248626, 248626, 248626, 248626, 248626, 248626, 248626, 248626, 248626, 248626, 248
						27486265, 35695763, 50431602, 10106370, 20281089, 264629, 18108374, 18108376,
						55811576, 35696423, 35695855, 264634, 264635, 264636, 264555, 60431850, 264556.
852	10147366 (1703, 1704)	170				264691

- 1					
٦	13032587 (1705, 1706)				264636
854	80052438 (1707, 1708)	80052438 (1707, 1708) Novel Protein sim. GBank gij3402836jemb[CAA/6082] - (Y16136) 2-enoate reduciase (Moorella thermoacetica)		reductase	204200
855	79641130 (1709, 1710)				264692
856	11594238 (1711, 1712)		,	UNCLASSIFIED	264591
857	79210165 (1713, 1714)			UNCLASSIFIED	264630, 264634
858	80248910 (1715, 1716)				265008, 265009, 264601, 264602, 264603. 18108351
950	20206634 (1717 1718)				284559
800	R0041749 (1719 1720)			UNCLASSIFIED	264489
861	85857045 (1721, 1722)			UNCLASSIFIED	33657023, 264630
862	80079467 (1723, 1724)				264600
863	80579931 (1725, 1726)	80579931 (1725, 1726) Novel Protein sim. GBank gij2246532 (U93872) - ORF 73.		UNCLASSIFIED	264488, 18108398, 35696286, 264259,
	•	contains large complex repeat CR 73 [Kaposi's sarcoma- associated herpesvirus]			18108351, 264288, 265021
864	94939904 (1727, 1728)			UNCLASSIFIED	264259, 264112, 263974
865	80045310 (1729, 1730)	Г	Contains protein domain (PF01479) -		284635, 264600, 284836, 264591, 264602,
	-	stein (Streptomyces coelicolor	S4 domain		264693
998	80162031 (1731, 1732)	Protein sim. GBank		transport	264288, 284557, 264558
		gi[4557876]ref[NP_000341.1[pABCR - ATP binding cassette transporter			
867	80062402 (1733, 1734)				264605
888	10075364 (1735, 1736)			UNCLASSIFIED	264909
869	80062406 (1737, 1738)				264605, 264687, 18108374
870	80249651 (1739, 1740	80249651 (1739, 1740) Novel Protein sim. GBank gil628660[pir][S37755 - Adenylyl- Iransferasa - Escharichia coli		transferase	264601, 264636
	20202000 11211	20230005 (4744 4740) Niveri Destriction in Contra		INCI ASSISIED	264603
<u>.</u>	Z03/8Z85 (1741, 1742 	gli1708180[sp[Q10602]HEMK_MYCTU - HEMK PROTEIN HOMOLOG		ONCLASSIFIED	500403
872	95197114 (1743, 1744	95197114 (1743, 1744) Novel Protein sim. GBank gil1545959lemblCAA67763  -		UNCLASSIFIED	35696286, 22278998, 264259, 29331822,
		(X99384) paladin [Mus musculus]			29331824, 29331825, 60432289, 29331826,
			-		29331627, 29331629, 33090032, 204309, 384608, 384608, 384607, 384608, 284619
					204903, 204900, 204907, 204900, 204909,
					204310, 203000, 204311, 204312, 203001,   204010, 20
_					Z03000, Z03003, Z04810, Z04331, 00433330,
					264596, 52646517, 67 166474, 265010,
					264602, 264603, 265017, 265018, 264605,
					10100331, 204704, 204700, 204700,   E2644230 264760 21006768 265021
					02644229, 204709, 21900709, 205021,   054534   054529,
					135696423, 65274791, 264631, 264632,
	-				284635, 264636, 264556, 264637, 264638,
					264639, 60432113, 22279000, 22279002,
1	22.54 24.54 005.00		3000000	TAICI ACCIETED	264506
8/3	20189728 (1745, 1746	20189728 (1745, 1746) Novel Protein sim. GBank gi[4156104 (AE001589) - putative   Outer membrane protein [Helicobacter ovlori J99]	gij4156104 (AE001569) - putative Contains protein domain (PF00096) - UNCLASSIFIED Helicobacter pylori J99)	UNCLASSIFIED	C8C407

264600	264259, 264448, 264288, 264557, 87168518	264907, 264601, 264602, 264605, 265020. 60431602	22278998, 264909, 264369	264369, 264555	56182575, 29331824, 264508, 264906, 265018, 18108351, 264448, 264683, 21906768, 21908768, 60170615, 33657023, 65274620, 33657109, 18108374, 35695855, 264563	264508	264508	204330	204080	050407	264605, 18108362	264690	35696052, 264906, 264600, 264003, 35695917, 35695855, 264636	264908	264603	264629
UNCLASSIFIED				ATPase_associated 264359, 264359		cathepsin	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED		dehydrogenase	UNCLASSIFIED		
		Contains protein domain (PF00005) - transport ABC transporter	Contains protein domain (PF00320) - transcriptfactor GATA zinc finger			Contains protein domain (PF00051) - cathepsin Kringle domain										
80077692 (1747, 1748) Novel Protein sim. GBank gij134319 sp P07819 SCRB_BACSU - SUCROSE-8-	GBank	ment) in sim. GBank gij3128283 (AF010496) - iron(iii) nsport ATP-binding protein (Rhodobacter	G7802548 (1753, 1754) Novel Protein sim. GBank gil731074spiP40349URB1_USTMA - SIDEROPHORE RIOSYNTHESIS REGIII ATORY PROTEIN URBS1	S) Novel Protein sim. GBank gij1351614[sp]Q09853[YAED_SCHPO - HYPOTHETICAL 18.1 6 KD PBOTEIN C2303 13C IN CHROMOSOME I	94328962 (1757, 1758) Novel Protein sim. CBank gil3875304[emb[CA589434] - (274639) predicted using Genefinder; CDNA EST EMBL:C07609 comes from this gene; cDNA EST EMBL:C09023 comes from this gene; cDNA EST yk489h9.3 comes from this gene; cDNA EST yk489h9.5 com			11077011 (1763, 1764) Novel Protein sim. GBank gilz632098 emb CAA75667  - (Y15513) Prodos protein [Drosophila melanogaster]		13517921 (1767, 1788) Novel Protein sim. GBank gij1155068jembjCAA6425j - (X94976) cell wall-plasma membrane linker protein IRrassica nanus	80052457 (1769, 1770) Novel Protein sim. GBank gil2078027 lemb CAB08467  - (295208) hypothetical protein Rv2372c [Mycobacterium	-	94315307 (1773, 1774) Novel Protein sim. GBank gij2695834jembjCAA15904j -	10083399 (1775, 1776) Novel Protein sim. GBank gij5689395 dbjjBAA82981.1 - ILARD28453 KIAA1029 orolein Homo sabiens	20385917 (1777, 1778) Novel Protein sim. GBank gil1881338 dbj BAA19365  - (AB001488) PROBABLE INTEGRAL MEMBRANE PROTEIN, SIMILAR TO CHLORAMPHENICOL RESISTANCE PROTEIN OF STREPTOMYCES	VENEZUELAE.   Bacillus subtilis    19904337 (1779, 1780)   Novel Protein sim. GBank gi 854065 emb CAA58337  -   (X83413) U88 [Human herpesvirus 6]
10077692 (1747, 1748)	(1749, 1750)	16465157 (1751, 1752)	87802548 (1753, 1754)	80187289 (1755, 1756)	94328962 (1757, 1758	8491135 (1759, 1760)	11290122 (1761, 1762)	11077011 (1763, 1764	79582969 (1765, 1766)	13517921 (1767, 176)	80052457 (1769, 177	11685138 (1771 1772)	94315307 (1773, 177	10083399 (1775, 177	20385917 (1777, 177	19904337 (1779, 178
874 8	875 8	876	877	878	879	980	<u>8</u>	882	88	884	885	ä	887	888	889	890

264636	(3030330 00070000 0000000	22278996, 22278999, 29331828, 35595022. 264908, 264909, 265009, 265011, 284602. 265019, 264766, 21906565, 21906766.	21906/09, 263020, 263021, 3032330		263978	264508, 264600, 264555, 264559	264632	264686, 29331828, 264511	56182575, 60432289, 56182435, 60432229,	55811957, 22279000, 264486	264601	264769, 264691, 264563	264907, 264602, 264605, 264769, 35695917.	18108376, 264563	264999, 265008, 284602, 264604, 264769, 264689, 264693	25505053 264005 284510 284511 284512	264605, 264760, 18108351, 284762, 284687,	264768, 264769, 264688, 21908764, 35695917, 27486262, 35695855, 264634,	264636, 264486	264604	264909	264595, 264605		60432289, 264601, 264690	264638	264602	265009
UNCLASSIFIED		transcriptfactor	oschious	Symnaso Minaso		UNCLASSIFIED		UNCLASSIFIED				transport	synthase		isomerase		Kinase			hydrolase	UNCLASSIFIED			UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	- eph
								Contains protein domain (PF00047) - UNCLASSIFIED Immunoglobulin domain					Contains profein domain (PE01411) - synthase	IRNA synthetases dass II (A)	Contains protein domain (PF00330) - isomerase Aconitase family (aconitate	nydratase)	Contains protein domain (PF00370) - kinase FGGY family of carbohydrate	kinases		Contains protein domain (PF00491) - hydrolase							Contains protein domain (PF00684) - eph DnaJ central domain (4 repeats)
1	gi[4959396 gb AAD34331.1 AF11248 - (AF112481) RAD54B  protein [Homo sapiens]	87634157 (1783, 1784) Novel Protein sim. GBank gil545526 bbs 143833 - LBP- Partanscription factor binding to initiation site of HIV-1 (allematively spliced) [human, Namawa cells, Peptide, 541	a)	Novel Protein sim. GBank gij2829688jspjP80608jCYSK_MAIZE - CYSTEINE SYNTHASE (O-ACETYLSERINE SULFHYDRYLASE) (O- ACETYLSERINE (THIOL)-LYASE) (CSASE)								Vovel Protein sim. GBank gi 1750127 (U66480) - YncC	Bacillus subtilis	Novel Protein sim. Gbank gij3122879[sp]007438[SYA_MYCTU - ALANYL-TRNA SYNTHFTASF (ALANINE-TRNA LIGASE) (ALARS)	80503301 (1803, 1804) Novel Protein sim. GBank gij3355701[emb CAA20001] - (AL031124) 3-isopropyimalate dehydratase large subunit	Streptomyces coelicolor	82060208 (1805, 1806) Novel Protein sim. GBank gil2960120 emb CAA18018.1  -			20451078 (1807, 1808) Novel Protein sim. GBank	gif Econ 1-pt Novel Protein sim. GBank gi4567200[gb]AAD23616.1[AC00716 - (AC007168)	hypothetical protein (Arabidopsis thailana)	80052626 (1611, 1612) Novel Frotein sim. Graans 915/56200jennjc/Avz.1252  (ALO31855) putative vacuolar membrane protein [Schingsaccharomyces numbel			20727907 (1817, 1818) Novel Protein sim. GBank gij3868940 dbj BAA34296  -	16776206 (1819, 1820) Novel Protein sim. GBank gil4589726 dbj BAA76883.1] - (AB003137) DnaJ homolog protein {Salix gilglana}
13516879 (1781, 1782) Novel Protein sim. GBank	<u> </u>	87634157 (1783, 1784) NO	aa	78168037 (1785, 1786) Novel Protein sim. GBank gij2829688[sp P80608]CY- SYNTHASE (O-ACETYLS ACETYLSERINE (THIOL)	14402340 (4787 4788)	11102240 (1/6/, 1/60)	70777903 (1703, 1793)	94991923 (1793, 1794)	87805100 (1795, 1795)	(0011, 0011) 80108010	11100463 (1797 1798)	4=-		80502410 (1801, 1802) Novel Pfotein sim. GBank gij3122879 sp 007438 SY SYNTHETASE (ALANINE	80503301 (1803, 1804) h		82060208 (1805, 1806)			20451078 (1807, 1808)	9398483 (1809, 1810)	100000000000000000000000000000000000000	80052626 (1811, 1612)	_	11754482 (1815 1818)	20727907 (1817, 1818)	16776206 (1819, 1820)
891		892		893	Т	Т	C S		808	020	000	88		100	902	_	803			8	905		<u>§</u>	00 20	Š	8	910

911	87454340 (1821, 1822)	87454340 (1821, 1822) Novel Protein sim. GBank gi[548774[sp P35685 RL7A_ORYSA - 60S R!BOSOMAL PROTEIN L7A		prof	265010, 264604, 60432113
912	20448863 (1823, 1824)	20448863 (1823, 1824) Novel Protein sim. GBank gi[2314008 gb AAD07921.1  - (AE000597) CDP-diglyceride hydrolase (cdh) [Helicobacter pylori 26695]			264559
913	20469357 (1825, 1826)			IFIED	264604
914	79183351 (1827, 1828)	Novel Protein sim. GBank gild 17657ispjQ03604 RIR1_CAEEL - PROBABLE RIBONUCLEOSIDE-DIPHOSPHATE REDUCTASE LARGE CHAIN (RIBONUCLEOTIDE REDUCTASE)	Contains protein domain (PF00317) - reductase Ribonucleolide reductase		264636
915	87606703 (1829, 1830)	87606703 (1829, 1830) Novel Protein sim. GBank gil5689571(db) BAA83069.1(- (AB029040) KIAA1117 protein [Homo saplens]			18108398, 22278996, 66714117, 264906, 264591, 21906768, 265020, 55811576, 264638
916	79444091 (1831, 1832)	79444091 (1831, 1832) Novel Protein sim. GBank gil4188110jemb CAA71790j - (Y10831) putative integrase [Ralstonia eutropha]			264595
917	20195985 (1833, 1834)			SIFIED	264605
918	91226795 (1835, 1836)	91228795 (1835, 1836) Novel Protein sim. GBank gil1655699 emb CAA69032  - (Y07752) pherophorin-S [Volvox carteri]		synthase	264259, 29331826, 264908, 265019, 264448, 265020, 264635, 83373044
919	80436785 (1837, 1838)	80436785 (1837, 1838) Novel Protein sim. GBank gil5689968 emb CAB52005.1 - (AL109663) putative membrane protein [Streptomyces			265006, 264512, 264600, 264602, 264604, 264768, 18108370, 264563
920	79606095 (1839, 1840)	78606095 (1839, 1840) Novel Protein sim, GBank		peptidase	264508
		(9) (1168448 sp)(205813 AMP1_STRLI - XAA-PRO gill 168448 sp)(205813 AMP1_STRLI - XAA-PRO gaminopeptidase (X-PRO aminopeptidase I) (AMINOACYLPROLINE AMINOPEPTIDASE I)	·		
921	19858634 (1841, 1842)	19858834 (1841, 1842) Novel Protein sim. GBank gij3850084jemb CAA21911.1 - (AL033389) alcohol dehydrogenase [Schizosaccharomyces		UNCLASSIFIED	264600
629	78982605 (1843 1844)				265019, 22279002
923	86695830 (1845, 1846	86695830 (1845, 1846) Novel Protein sim. GBank gi[267079 sp P29514 TBB6_ARATH - TUBULIN BETA-6		tubulin	264907, 265008, 265009, 265010, 18108351, 264689, 265021, 18108370, 18108374, 18108385
924	21431341 (1847, 1848)				264510
928	20630332 (1849, 1850	20630332 (1849, 1850) Novel Protein sim GBank		esterase	264603
		git497688 spig660963 PAFA_MOUSE - PLATELET- ACTIVATING FACTOR ACETYLHYDROLASE DEBECT DOOR JAKE ACETYL HYDROLASE - JOHE 2			
		ACYLHYDROLASE) (LDL-ASSOCIATED			
		PHOSPHOLIPASE A2) (LDL-PLA(2)) (2-ACETYL-1- ALKYLGLYCEROPHOSPHOCHOLINE ESTERASE) (1- ALKYL 2-ACETYLCI VCEROPHO			·
926		79397657 (1851, 1852) Novel Protein sim. GBank gil3882325 dbj BAA34522.1 - (AB018345) KIAA0802 protein IHomo saplens)			558   1957, 263972, 264639
827	37036201 (1853, 1854)			UNCLASSIFIED	264769

828	80070610 (1855, 1856)	80070610 (1855, 1856) Novel Protein sim. GBank gil 156146 (M30316) - xanthine		dehydrogenase	264605
╗		dehydrogenase (AA al 2538) (Calliphora vicina)		I MICH ACCIDION	264803
				UNCLASSIFIED	204003
930	5496348 (1859, 1860) N				264259
931	10245731 (1861, 1862)	4490609 emb CAB38642.1  - eductase major subunit	Contains protein domain (PF00317) - reductase Ribonucleotide reductase	reductase	264486
932	80420613 (1863, 1864)	80420613 (1863, 1864) Novel Protein sim. GBank gil5459396 emb CAB50754.11- (AL096839) putative integral membrane transport protein  Streptomyces coelicolor		transport	264508, 264905, 264906, 264909, 264600. 264602, 264603, 264605, 264766, 264558, 18108387, 264486
933	94326010 (1865, 1866)		Contains protein domain (PF00627) - UNCLASSIFIED UBA domain	UNCLASSIFIED	264508, 264686, 264693, 27486261, 18108370, 65274791, 264636, 264559, 22279002
934	80039105 (1867, 1868)	80039105 (1867, 1868) Novel Protein sim. GBank gil119111 sp P12978 EBN2_EBV - EBNA-2 NUCLEAR PROTEIN			264369, 22279002
935	80063162 (1869, 1870)			UNCLASSIFIED	264604, 264605, 264693, 18108370, 18108374
936	80026632 (1871, 1872)	80026632 (1871, 1872) Novel Protein sim. GBank gij845686 (M32103) - ORF-27  Staphylococcus aureus		UNCLASSIFIED	22278995, 22278996, 264602, 264687, 32833986, 18108387
937	80250273 (1873, 1874)	80250273 (1873, 1874) Novel Protein sim. GBank gil1360669ipirIICGHU1V - collagen alpha 1(V) chain precursor - human	٠	synthase	263978
938	80026633 (1875, 1876)	80026633 (1875, 1876) Novel Protein sim. GBank gi 2226243 emb CAA74531.1  -  (Y14083) hypothetical protein [Bacillus subtilis]		UNCLASSIFIED	264602
939	11071694 (1877, 1878)	-			264600
940	94144252 (1879, 1880)	94144252 (1879, 1880) Novel Protein sim. GBank gij3560166jemb CAA20678j -	Contains protein domain (PF00442) - ubiquitin	ubiquitin	264905, 264906, 264907, 264908, 264909. 264511, 264910, 264592, 33657402, 264596,
		(Schizosaccharomyces pombe)	hydrolases family 2		264758, 264760, 264683, 264768, 264768, 264769, 23657023, 33657109, 264628, 264629, 264630, 264636, 264555, 264637, 264638, 264638, 83373044
					18108385, 264565, 18108391
2 2	11398414 (1881, 1882)			I INCLASSIFIED	264760
2 63	80080258 (1885, 1886)	80080258 (1885, 1886) Novel Protein sim. GBank gil4033729 (AF038595) -		UNCLASSIFIED	264600, 264687, 264689, 264563
	,				
8	80216096 (1887, 1888)	80216096 (1887, 1888) Novel Protein sim. GBank		synthase	264511, 264603
		gi 2494764 sp Q50729 GUAA_MYCTU - GMP SYNTHASE (GLUTAMINE-HYDROLYZING) (GLUTAMINE AMIDOTRANSFERASE) (GMP SYNTHETASE)			-
845 5	80052477 (1889, 1890)	80052477 (1889, 1890) Novel Protein sim. GBank gij732353 spjP39606 YWCH_BACSU - HYPOTHETICAL 36 6 KD PROTEIN IN OOXD-VPR INTERGENIC REGION	Contains protein domain (PF00818) - UNCLASSIFIED Ice nucleation protein repeat	UNCLASSIFIED	264906, 264604, 264605, 265020, 18108387
946	79248402 (1891, 1892)	_			265017

osphatase		inscriptfactor		IFIED						٦			UNCLASSIFIED 264259	263978
Contains protein domain (PF00459) - phosphatase Inositol monophosphatase family	jū.	Contains protein domain (PF00096) - transcriptfactor 5 Zinc finger, C2H2 type	Contains protein domain (PF00038) - struct Intermediate filament proteins	î		tra (tra	j qe	Contains protein domain (PF00076) - dna_rna_bind 9.1[pSIP] - SYT interacting protein RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)		<b>D</b>		3		
	88165538 (1895, 1896) Novel Protein sim. GBank gil2827284 (AF041037) - novel antagonist of FGF signaling [Homo sapiens]	88081786 (1897, 1898) Novel Protein sim. GBank gil4507985 ref NP_003427.1 p2NF1 - zinc finger protein 135 Zinc finger, C2H2 type (clone pHZ-17)	78485872 (1899, 1900) Novel Protein sim. GBank gil1079461[pirt]S43865 - [cytokeratin 8, type II - potoroo (fragment)	20451411 (1901, 1902) Novel Protein sim. GBank gij5420387[emb CAB46679.1] - [AJ243459) proteophosphoglycan [Leishmania major]	Novel Protein sim. GBank gij5305702jgbjAAD41779.1jAF12686 - (AF126867) calpain- like protease (Mus musculus)	10196003 (1905, 1906) Novel Protein sim. GBank gij2495642 sp Q47142 YFHS_ECOLI - HYPOTHETICAL 41.8 KD PROTEIN IN CSIE-GLYA INTERGENIC REGION	Novel Protein sim. GBank gil2360965 (AF016253) - D- amino acid dehydrogenase [Klebsiella aerogenes]	95313410 (1909, 1910) Novel Protein sim. GBank gil5454064 ref NP_006319.1 pSIP  - SYT interacting proteir	80064224 (1911, 1912) Novel Protein sim. GBank gij2052129jembjCAB08155j - (294752) rimJ [Mycobacterium tuberculosis]		80036448 (1915, 1916) Novel Protein sim. GBank gil1709767!splQ00451 PRF1_LYCES - 36.4 KD PROLINE. RICH PROTEIN	80026647 (1917, 1918) Novel Protein sim. GBank gi[2131050jemb CAB09260j - (295844) opcA [Mycobacterlum tuberculosis]	37815406 (1919, 1920) Novel Protein sim. GBank gi[2129478 pir  S51939 - chitinase (EC 3.2.1.14) precursor - beet	
81802699 (1893, 1894) Novel Protein sim. GBank (AL021899) hypothetical p tuberculosis]	88165538 (1895, 1898) <sup>B</sup>	88081786 (1897, 1898)   	79485872 (1899, 1900) I	20451411 (1901, 1902)	79566954 (1903, 1904)	10196003 (1805, 1906)	9893326 (1907, 1908)	95313410 (1909, 1910)	80064224 (1911, 1912)	80056206 (1913, 1914)	80036446 (1915, 1916)	80026647 (1917, 1918)	37815406 (1919, 1920)	20567383 (1921, 1922)

264510, 264288, 264555, 264556, 264559, 264486	264112, 264910, 264689	<u>264488, 265017, 264448, 264634, 264558, </u>	264906, 264592, 264596, 264604, 264768,	21906764, 264692, 264693, 264629, 264636, 264638	264760	264693	35696286, 264685, 264686, 35695917, 264692, 18108374, 264635	2278996, 22278998, 22278999, 29147620, 264828, 255006, 265007, 265008, 265009, 48108348, 33109954, 265010, 265011,	18108351, 264288, 21906767, 21906768, 18108370, 18108374, 18108377, 264630, 264635, 18108380, 83373044, 18108387, 18108388	265007, 265020, 22279002	264557	264604	264565	26451U	[264508, 264808, 264739, 26403£, 26408;	264353	264906	ACTION OF SERVICE SERVICES	264600, 264602, 264603, 264604	284767, 284768, 265006, 265007, 204500	204000, 204002, 204003 20,200 204656 204658 204547 264559	264692, 204333, 204330, 204301, 20130
		UNCLASSIFIED	I INCI ASSIFIED				dna_ma_bind	UNCLASSIFIED		UNCLASSIFIED			UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	,	UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED
					Contains protein domain (PF00072) - phosphatase Response regulator receiver domain		Contains protein domain (PF00096) - dna_ma_bind Zinc finger, C2H2 type															
	78832019 (1927, 1928) Novel Protein sim. GBank gil4589622 dbj BAA76833.1 -	(AB023206) KIAA0989 protein Homo sapiens)	(A.1243459) proteophosphoglycan (Leishmania major)		79255708 (1933, 1934) Novel Protein sim. GBank gij1731207[sp]Q11156[RGX3_MYCTU - SENSORY TEANSON CONTINENT PROTEIN REGX3	79560269 (1935, 1936) Novel Protein sim. GBank gilz661836 emb CAA75187  - (Y14964) putative transport protein [Methylophilus	metnylotrophus 79919470 (1937, 1938) Novel Protein sin. GBank gi[5419878 emb CAB46422.1 - /Ast 094747 hymothetical protein (Homo sapiens)					20370183 (1945, 1946) Novel Protein slm. GBank 20370183 (1945, 1946) Novel Protein slm. GBank gilj1723119[spp.53990]Y174_HUMAN - HYPOTHETICAL	PROTEIN KINAUTA		80205742 (1951, 1952) Novel Protein slm. GBank gil3881459 emb CAA92988.1  -	(268753) predicted using Genefinder, Similarity to Yeast hypothetical protein YIK9 (SW:YIK9_YEAST); cDNA EST EMBL:D27680 comes from this gene; cDNA EST EMBL:D27679 comes from this gene; cDNA EST	10355349 (1953 1954) Novel Protein sim. GBank	gissesispidos335iXYS3_PSEPU - XYLDLEGF OPERON TRANSCRIPTIONAL ACTIVATOR 3		Novel Protein sim. GBank gij3171904(emb CAA75869  - (r/15908) DIA-12C protein [Homo saplens]		Novel Protein sim. GBank gi[3599940 (AF017368) -
80590374 (1925, 1926)	9832019 (1927, 1928) N	() N (0264 1930) N	()	95292815 (1931, 1932)	9255708 (1933, 1934) N	19560269 (1935, 1936) h	19919470 (1937, 1938)	95085947 (1939, 1940)		10000 1001 001000	78919770 (1941, 1942)	20710/04 (1943, 1944) 20370183 (1945, 1946)	80057103 (1947 1948)	10196018 (1949, 1950)	80205742 (1951, 1952)		10355349 (1953 1954)	the section of the se	R0025927 (1955 1956)	80447820 (1957, 1958)	80025928 (1959, 1960)	80098550 (1961, 1962)
963 80	964	985		996	7 298	9968 7	696	970		┪	Т	972	72	Т	978		077		07R	979	980	8

982 6	80195670 (1963, 1964) N	80195670 (1963, 1964) Novel Protein sim. GBank gil2950220jemb CAA71575 -		UNCLASSIFIED	264404
+	)	(Y10545) fused-ccdB [Eschenchia coli]		struct	65274572, 56182575, 264908, 264909.
	90995041 (1965, 1966) N	90995041 (1965, 1966) Novel Protein sim. Gbank gil476363 ptill043402 - tilyosiit Theavy chain-B neuronal - chicken			265007, 265008, 264758, 265010, 55811150,
_	<u>-</u>				33657023, 284634, 264557, 284558
†	20466876 (1967, 1968)		1		264605
+=		im. GBank gij3451504jembjCAA07660.1j - bothetical protein BbLPS1.21 [Bordetella	Contains protein domain (PF00534) - transferase Glycosyl transferases group 1		56182435, 264600
7	_	Oroncuisepuca		UNCLASSIFIED	264106, 264110, 265020, 60170615
1	3080102808 (1871, 1872)			Γ	264909
1	1900/231 (19/3, 19/4)				264600
000	19030001 (1973, 1970)			UNCLASSIFIED	264508, 265017, 264534, 264564
Т		MEDOCAE ARCCOACO	Contains protein domain (PF01421) - oxidase	oxidase	264259, 264908, 265009, 264910, 264596, 264369, 264288, 264768, 264628, 264635,
					264568
T	10106140 (1981 1982)				264909
266	79845694 (1983, 1984)	79845694 (1983, 1984) Novel Protein sim. GBank gil2105049jembjCAB08835j - (285436) hypothetical protein Rv3645 [Mycobacterlum	Contains protein domain (PF00211) - UNCLASSIFIED Adenylate and Guanylate cyclase	UNCLASSIFIED	264508, 264593
Т	-	tubercuiosis			264907
222	10014033 (1803, 1800)				264603
	11090590 (1987, 1988)	11090590 (1987, 1988) Novel Protein sim. GBank gij3329297 (AE001355) - Ribonudeoslde Reduclase, Large Chain [Chlamydia Irachomatis]		reductase	700607
200	04321011 (1080 1000)	04224044 (4080 1000) Novel Protein eim Chank	Contains protein domain (PF00176) -	helicase	18108398, 65274572, 22278996, 264490.
1	843Z1911 (1969, 1990)		SNF2 and others N-terminal domain		60432049, 29331827, 29146498, 264508, 264905, 264908, 56182435, 265008, 264591, 264592, 60431229, 60431735, 33657402, 264595, 264758, 21906754, 265010, 265017, 265018, 264605, 264766, 21906768, 21906769, 25811957, 264632, 264629, 264629, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264569, 264568, 203333044, 222739002, 264568, 55811957,
986	91013745 (1991, 1992)	91013745 (1991, 1992) Novei Protein sim. GBank gi 2911719 (AC00422/) -  KIA001LB [Homo sapiens]	Contains protein donain (Froussy) - prospiratese PDZ domain (Also known as DHR or GLGF).	per pudental	56526486, 87168518, 284910, 264906. 264565, 264566, 264693, 264766
266	80503347 (1993, 1994)	80503347 (1993, 1994) Novel Protein sim. GBank gij2649101 (AE001001) - ABC transporter, ATP-binding protein (Archaeoglobus fulgidus)	Contains protein domain (PF00005) - transport ABC transporter	· transport	35696286, 22278897, 22278999, 204509, 264905, 264908, 265010, 264600, 264602, 264605, 264688, 264769, 265021, 264565, 18108391
866	11397390 (1995, 1896)	11397390 (1995, 1996) Novel Protein sim. GBank gij123530lspjP04929JHRPX_PLALO - HISTIDINE-RICH GLYCOPROTEIN PRECURSOR		,	264595

				ſ	
666	11768047 (1997, 1998)	11768047 (1997, 1998) Novel Protein sim. GBank		UNCLASSIFIED	264682
		gijz506697 sp P46490 YFCA_HAEIN - HYPOTHETICAL  PROTEIN HI0198			
1000	20727944 (1999, 2000)			UNCLASSIFIED	264602
T	86673131 (2001, 2002)	86673131 (2001, 2002) Novel Protein sim. GBank gil2224699 dbj BAA20833  - (AB002377) KIAA0379 [Homo sapiens]	Contains protein domain (PF00023) - kinase Ank repeat		60432049, 264907, 264909, 264511, 264603, 264683, 264684, 264687, 264689, 29146627, 21906769, 264692, 18108385, 22279000
1002	80189603 (2003, 2004)	80189603 (2003, 2004) Novel Protein sim. GBank ai15861211spP37709ITRHY RABIT - TRICHOHYALIN		struct	265009, 264369, 265020
1003	17833491 (2005, 2006)				265019
	16314987 (2007, 2008)	16314987 (2007, 2008) Novel Protein sim. GBank gil854065 emb CAA58337  - (X83413) U88 IHuman herpesvirus 6]	•		264635
1005	79617144 (2009, 2010)	79617144 (2009, 2010) Novel Protein sim. GBank		UNCLASSIFIED	264508
		gij114073jspjP07672JAPT_ECOLI - ADENINE PHOSPHORIBOSYLTRANSFERASE (APRT)			
1006	37815429 (2011, 2012)			UNCLASSIFIED	264259
_	79620871 (2013, 2014)	+=-		synthase	264905
1008	88094444 (2015, 2016)	88094444 (2015, 2016) Novel Protein sim. GBank gij2808807 emb CAA04607.11 -		synthase	265007, 264602, 264605, 264760, 264536
		(AJ001206) putative trehalose synthase [Streptomyces coelicolor]			
1009	57451289 (2017, 2018)	57451289 (2017, 2018) Novel Protein sim. GBank gij3639077 (AF090113) - AMPA	Contains protein domain (PF00595) - kinase	kinase	264102, 264288
		receptor binding protein [Rattus norvegicus]	PDZ domain (Also known as DHR or GLGF).		
1010	94672537 (2019, 2020)		a	dehydrogenase	264592
		NADH-dependent oxidase, may function as a demethylase [Sinorhizoblum melitoti]			
1011	85546916 (2021, 2022)	85546916 (2021, 2022) Novel Protein sim. GBank gi 2342647 gb AAB86591.1  -		UNCLASSIFIED	35696052, 264905, 264764, 264768,
	-	(U90653) DHHC-domain-containing cysteine-rich protein [Homo sapiens]			35695917, 264629
1012	95294456 (2023, 2024)	95294456 (2023, 2024) Novel Protein sim. GBank gij3413411 jemb CAA20272] -	Contains protein domain (PF00013) - phosphorylase	phosphorylase	35696052, 264905, 264600, 264601, 264602,
		(AL031231) guanosine pentaphosphate synthetase/ polyribonucleotide nucleotidyltransferase (Streptomyces	KH domain		264605, 264762, 264769, 264768, 264669
		coelicolor			
1013	86095772 (2025, 2026)			UNCLASSIFIED	264591, 21906/68
1014	1014 86608828 (2027, 2028)				29331824, 265019, 265020

264600	3027505	204393	265010, 265011, 264600, 264602, 264603,	264605, 264769, 18108372, 18108374	264905, 264906, 264909, 264595, 264692, 264630, 264634, 264638	264689	35696052, 264906, 264510, 18108354,	264687, 264769, 264689, 60431602, 18108385, 264486	264906	264692	264905, 66712502, 264908, 264766	29331824, 264909, 60433438, 265019	264604, 264634	264002 264003 264094 264683 264689.	263967	264595	264907		264605	264565, 264567	ACCADE 254503 355017	Z85006, Z8460Z, Z85011	28331823, 28403,	000700			264687		264905, 264634		264686
UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED		OSISIED AND A STREET	UNCLASSIFIED	isomerase	kinase			- transport		UNCLASSIFIED			helicase	UNCLASSIFIED	- UNCLASSIFIED		UNCLASSIFIED		UNCLASSIFIED
Contains protein domain (PF00330) - UNCLASSIFIED Aconitase family (aconitate	hydratase)		•										Contains protein domain (PF01477)	PLA I/LHZ domain		Contains protein domain (PF01131) - isomerase Prokaryotic DNA topolsomerase				Contains protein domain (PF00005) - transport	ABC transporter						Contains protein domain (PF00220) - UNCLASSIFIED				
1028   20297928 (2055, 2056) Novel Protein sim. GBank gilz791409[emb]CAA16003] - (4. 021184) acn fMycobacterium tuberculosis]						lovel Protein sim. GBank	gilagosagoillarine on the control of	Novel Frotein Siff. Spark glassess (Account) hypothetical protein (Aquifex aeolicus)	Novel Protein sim. GBank gil405895 (U00007) - methionyl-	אואר שלוווופוסטס לבשתכוומוום כמוו			Novel Protein sim. GBank gil790819 (L39891) - polycystic	kidney disease-associated protein (Homo sapiens)	Novel Protein sim. GBank gij2605967 (AF030027) - 24 Fruine hernesvirus 41	Novel Protein sim. GBank gilp352095 (U97022) - DNA	10156682 (2081, 2082) Novel Protein sim. GBank gij3256535(dbjjBAA29218.1) -	(AP000001) 301aa long hypothetical 2-phosphoglycerate kinasa (Pvrococus horisoshii)	Novel Protein sim. GBank gi 2058299 emb CAA66953 -	(X98309) ARI protein   Urosophila melanogaster  80057136 (2085, 2086)   Novel Protein sim, GBank gil1870167 emb CAA70125  -	(Y08921) msiK (Streptomyces reticuli)		52415482 (2089, 2090) Novel Protein sim. GBank gij5689890 emb CAB52053.1 -	(AE 1097.92) hypotitietical protein joinepromyces composed A3(2)]	11754862 (2091, 2092) Novel Protein sim. GBank gil854065 emb CAA58337  - (X83413) U38 (Human heroesvins 6)	37036258 (2093, 2094) Novel Protein sim. GBank gil4210471 dbj BAA74535.1  -		(ALUS1232) nypometical protein 3C 10n3.07 (Skeptoniyes) coelicolori	Novel Protein sim. GBank	gij5051636jgbjAAD38328.1jAF0/3/2 - (AF0/3/2/) EH Idomain-binding mitotic phosphoprotein [Homo saplens]	
0297928 (2055, 2056) NO		14665090 (2057, 2058)	1030 88095343 (2059, 2060)		95289117 (2061, 2062)	94673275 (2063, 2064) Novel Protein sim. GBank	6	86464818 (2055, 2065) N	79245937 (2067, 2068) Novel Protein sim. GBank	10705 0000 10000 0070)	79930333 (2009, 2070)	63604996 (2071, 2072) 63606066 (2073, 2074)		×	87260021 (2077, 2078) Novel Protein sim. GBank	80026840 (2079, 2080) Novel Protein slm. GBanl	10156682 (2081, 2082)		11084375 (2083, 2084) Novel Protein sim. GBanl	80057136 (2085, 2086)					_	37036258 (2093, 2094)	_		81755108 (2097, 2098) Novel Protein sim. GBar		1050 79471521 (2099, 2100)
1028		1029	080		1031	1032	_	1033	1034	300,	_	1030	1038		1039	1040	1041		1042	1043	<u>}</u>	19 4	<u>5</u>		1046	1047	1048		1049		1050

18108374, 264769, 265010, 265011, 284601, 265009, 264604, 264605, 264636, 18108351, 264692	264508, 264762, 264687, 264486	264686, 18108374, 29331824, 83373044, 21906754, 52845158, 56182435, 264689, 29331827, 21906765, 21906765, 21906765, 21906769, 25811957, 87168518, 256502, 265002, 265007, 265018, 22278900, 22278900, 22278900, 264482, 264906, 52644150, 264909, 264288, 29331822, 52645080, 264766	264686	264907, 264602, 264681, 264288, 21906768, 33657109, 55810764, 35695855, 264631	264906	264682	264764	35696052, 35695855, 265009, 264636	29146499, 264681, 264683, 264687	264634	264593	264508	265007, 265008, 18108351, 18108385
			SIFIED	helicase	mapolymerase	UNCLASSIFIED		transcriptfactor	glycoprotein	phosphatase	dehydrogenase		
	Contains protein domain (PF00389) - dehydrogenase D-Isomer specific 2-hydroxyacid dehydrogenases			Contains protein domain (PF00270) - helicase DEAD/DEAH box helicase									
	24_BACSU - D-3- EHYDROGENASE (PGDH)	94851640 (2105, 2106) Novel Protein sim. GBank gil5441319 emb CAB46717.1  - (AL034396) dJ1158B12.1 (zinc finger, X-linked, duplicaled A) [Homo sapiens]		80594138 (2109, 2110) Novel Protein sim. GBank gi 5052508 gb AAD38584.1 AF14560 - (AF145609) BCDNA,GH02833 [Drosophila melanogaster]	Novel Protein sim. GBank gij3021676[dbjjBAA25358] - (D86033) RNA polymerase sigma-70 factor [Pseudomonas fluorescens]	85657216 (2113, 2114) Novel Protein sim. GBank gi]1226281 (U50308) - No definition line found [Caenomabditis elegans]		94662754 (2117, 2118) Novel Protein sim. GBank gij1170016jspjP46808jGREA_MYCLE - TRANSCRIPTION ELONGATION FACTOR GREA (TRANSCRIPT CLEAVAGE FACTOR GREA)	79481169 (2119, 2120) Novel Protein sim. GBank gi[2499087]sp[009332]UGGG_DROME - UDP- GLUCOSE:GLYCOPROTEIN GLUCOSYLTRANSFERASE PRECURSOR (DUGT)	11034025 (2121, 2122) Novel Protein sim. GBank gij90254 pir  A28334 - protein- tyrosine-phosphatase (EC 3.1.3.48) Ly-5 precursor (B-cell variant) - mouse	39567937 (2123, 2124) Novel Protein sim. GBank gij3334200jspjO49954jGCSP_SOLTU - GLYCINE DEHYDROGENASE (DECARBOXYLATING) PRECURSOR (GLYCINE DECARBOXYLASE) (GLYCINE CLEAVAGE SYSTEM P-PROTEIN)	Novel Protein sim. GBank gi[2499986]sp[Q41228]PSE1_NICSY - PHOTOSYSTEM I REACTION CENTRE SUBUNIT IV A PRECURSOR (PSI-E A)	78891783 (2127, 2128) Novel Protein sim. GBank gij82654 pir  JA0086 - 10K zein precursor - malze
80475471 (2101, 2102)	82442962 (2103, 2104)	94851640 (2105, 2106)	79580225 (2107, 2108)	80594138 (2109, 2110)	17882319 (2111, 2112)	85667216 (2113, 2114)	80376576 (2115, 2116)			11034025 (2121, 2122)	39567937 (2123, 2124)	8490481 (2125, 2126)	78891783 (2127, 2128)
1051	1052	1053	1054		1056	1057	1058		1060	1061	1062	1063	1064

264600, 264602, 264689	900			264909	688, 18108362, 264558, 264600, 264/6U		264604	264604, 264760		264602	264687, 264688, 21906784, 35696052,	35695917, 3569585, 264600, 264601,	264690, 264906, 264762, 264628, 264768	264909	264906		56994075, 22278998, 264594, 264757.	264596, 265018, 265019, 264681, 264369. 264688, 265020, 18108364, 18108374	4769		264905		264600	8108394, 264769, 264634, 264636		264684	83373044, 265019, 22278002, 264482.	18108351, 264682, 264908, 264693, 264487	264758, 264768, 264769, 21806767, 204311, 264910, 264636, 264635, 264905, 264636.	264906, 264637, 264907, 264908, 264764, 264638, 20281099, 264766, 264595
transferase 2646	265000	skuinase					UNCLASSIFIED 264	$\neg$		synthase 264	INCLASSIFIED 264		7.00	26		,	UNCLASSIFIED 56	26	ATPase associated 264769	1						UNCLASSIFIED 2			UNCLASSIFIED 2	7 2
					Contains protein domain (PF00248) - reductase	Aldo/keto reductase family									Several (BODOOR) - levelthase	Contains protein contain (* 1 0000) FGE-like domain				,					-					
20021208 2120 2130   Novel Protein sim. GBank gi[2120998 pir] S70682 -			gilz506362 sp[P15042 DNLJ_ECOLI - DNA LIGASE	T	4007660lemhICAA223551 -	ein sim. Gbaink gilffoot oobjering ook ook ook ook ook ook ook ook ook oo	coelicolori	83002954 (2137, 2138) Novel Protein sim. GBank 914389484[dol] 547, 2.11 - (AB023143) KIAA0926 protein [Homo saptens]	82101992 (2139, 2140) Novel Protein sim. GBank gij120304 sp P15932 FLGK_SALTY - FLAGELLAR HOOK-	20710589 (2141, 2142) Novel Protein sim. GBank gil1750397 (U81261) - glutamate	synthase large subunit (Pseudomonas aeruginosa)					80105992 (2147, 2148) Novel Protein sim. GBank gil477532 pir  A49175 - Motch B	protein - mouse (fragment)	81850293 (2149, 2150) Nover Protein sim. Gearix Bisoso to Jeningson co. (Y17820) CALO protein (Drosophila melanogaster)		80477264 (2151, 2152) Novel Protein sim. GBank gij1176203 sp P46442 YHCM_ECOLI - HYPOTHETICAL 43.1 KD PROTEIN IN RPLM-HHOA INTERGENIC REGION	(F375)	79831334 (2153, 2154) Novel Protein sim. GBank gij4033487jspjQ44472JTUD4_AGRVI - PUTATIVE	HYDROXYPYROVAIE REDUCINSE	20288874 (2155, 2156)	)) Novel Protein sim. GBank gij3413628jembj-AAZU299 - (AL031260) hypothetical protein SC9A10.09 [Streptomyces	coelicolor		[(2	(4)	
021208 (2120 2130) N	. (	896879 (2131, 2132) IN	<u> </u>	_	10132178 (2133, 2134)	2062057 (2135, 2136)    (		3002954 (2137, 2138)	2101992 (2139, 2140)	0710589 (2141, 2142)		82356540 (2143, 2144)			79814400 (2145, 2146)	30105992 (2147, 2148)		81850293 (2149, 2150)		80477264 (2151 <b>,</b> 2152) '		79831334 (2153, 2154)		20288874 (2155, 2156,	80494518 (2157, 2158	•	11767188 (2159, 2160)	94747080 (2161, 2162)	81490656 (2163, 2164)	
00 300		1066 17		$\neg$	$\neg$	1088   82	_	1069 8:	1070 8:	1071		1072 8		_	1073 7		_	1075		1076		1077		1078	1079		1080	1081	1082	

60424179, 264905, 264908, 264510,	60432229, 264759, 87168474, 264005, 264769, 264689, 18108364, 18108376, 35695855, 264636	264769	60432229, 264687	264486	29331827, 264693	264905, 264601, 18108387	264488, 264511, 265011, 264682, 264768, 264689, 21906764, 35695917, 265020, 32833986, 18108370, 35695855	264604	264637	264566	264769, 35695917, 35695855, 254500, 264602, 264603, 264605, 18108351	264686	264906, 265007, 264595, 264600, 204602, 264603, 264604, 264605, 264762, 264766, 264769, 264636, 264558, 18108387, 60432113, 264482, 264486	264600, 264602, 264603, 264504, 264605, 35695917, 264692, 264631	265019	264687
UNCLASSIFIED		UNCLASSIFIED							UNCLASSIFIED			SIFIED	- nuclease	synthase	UNCLASSIFIED	synthase
			Contains protein domain (PF00118) - eph TCP-1/cpn60 chaperonin family	Contains protein domain (PF00238) - inbosomalprot Ribosomal protein L14			Contains protein domain (PF00329) - UNCLASSIFIED Respiratory-chain NADH dehydrogenase, 30 Kd subunit				Contains protein domain (PF00730) - nuclease Endonuclease III		Contains protein domain (PF00455) - nuclease Bacterial regulatory proteins, deoR family			
ovel Protein sim. GBank	gij 722845 spja 10523 jyo 1 n. myctu - hypothetical 44.6 kd protein cy427.23	Novel Protein sim. GBank gij418384jspjP32057jWCAI_ECOLI - PUTATIVE COLANIC ACID BIOSYNTHESIS GLYCOSYL TRANSFERASE WCAI	i 1899190 (U90204) - heat ella tvrosinosolvens	4_ACYKS - 50S RIBOSOMAL	sim. GBank gil4160198jemb CAA15431  - 327J16.3 (novel CHROMObox family protein)	Novel Protein sim. GBank gij2983155 (AE000693) - phosphoglucomulase/phosphomannomulase (Aquifex applicas)	1pE00177 - (AE001778) NADH ubunit, putative [Thermotoga	Novel Protein sim. GBank gil4007680 emb CAA22366  - (ALO34443) putative oxidoreduciase (Streptomyces coelicolori	12917471 (2181, 2182) Novel Protein sim. GBank gi[2495562]sp P77239 YLCD_ECOLI - HYPOTHETICAL 443 KD PROTEIN IN NFRB-PHEP INTERGENIC REGION PRECIESOR		80496304 (2185, 2188) Novel Protein sim. GBank gi[2960098 emb CAA17996.1] - (AL022121) nth [Mycobacterfum tuberculosis]	Novel Protein sim. GBank gil1001642 jdbijBAA10373  - (D64002) dGTP triphosphohydrolase [Synechocystis sp.]	87457250 (2189, 2190) Novel Protein sim. GBank gil4585587 emb CAB40855.1 - (AL049628) putative adenine glycosylase (Streptomyces coelicolor]	80025977 (2191, 2192) Novel Protein sim. GBank gij115001 sp P19206 BIOB_BACSH - BIOTIN SYNTHASE (BIOTIN SYNTHETASE)	-	79186424 (2195, 2198) Novel Protein sim. GBank gi[114135 sp P08205 ARGA_ECOLI - AMINO-ACID ACETYLTRANSFERASE (N-ACETYLGLUTAMATE SYNTHASE) (AGS)
1083 187446717 (2165 2166) Novel Protein sim. GBank	199	37799306 (2167, 2168) Novel Protein sim. GBank gil418384 sp P32057 WC/ ACID BIOSYNTHESIS GL	86475368 (2169, 2170) Novel Protein sim. GBank g	79608269 (2171, 2172) Novel Protein sim. GBank gil11729561splP46176IRL PROTEIN 14	79603979 (2173, 2174) N (A	79854963 (2175, 2176) Novel Protein sim. GBank phosphoglucomutase/phos applicated	80216800 (2177, 2178) Novel Protein sim. GBank gil4881768[gb]AAD36290. gehydrogenase, 30 kDa si marilima	11083825 (2179, 2180) Novel Protein sim. GBank (AL034443) putative oxido	12917471 (2181, 2182) N	80252286 (2183, 2184)	_	10880972 (2187, 2188)	87457250 (2189, 2190)	80025977 (2191, 2192)	79239560 (2193, 2194)	79186424 (2195, 2196)
1083		1084	1085	1086	1087	1088	1089	1090	1091	1082	- - - - - - - - - - - - - - - - - - -	5 4 8	1095	1098	1097	1098

264603	264259, 264636	264769	264910, 264909	264757	29331822, 21906754, 264555, 264556, 264558, 22279002.	264566	264555, 264369	264908, 264769	264907, 264510, 264511, 264600, 264602, 264605, 264768, 264769	264602, 264605, 264636	264908, 264593, 265010, 264601, 264603. 264604, 264605, 264682, 264769, 264693. 264636	18108370, 264557	264908	264600, 264602, 264604, 264605, 264762, 264769, 264565	264636	264686	26490R	201300
UNCLASSIFIED	desaturase	isomerase	UNCLASSIFIED	struct	UNCLASSIFIED		UNCLASSIFIED		transport	helicase				UNCLASSIFIED	dehydrogenase	INCI ASSIFIED	2000	
		Contains protein domain (PF01396) - isomerase Topoisomerase DNA binding C4 zinc finger																
Novel Protein sim. GBank gij3915144 spjO33017 TRMD_MYCLE - TRNA (GUANINE- N1)-METHYLTRANSFERASE (M1G- METHYLTRANSFERASE) (TRNA [GM37]	Novel Protein sim. GBank gij3023255ispjQ64420jACOD_MESAU - ACYL-COA DESATURASE (STEAROYL-COA DESATURASE) (FATTY ACID DESATURASE) (DELTA(9)-DESATURASE)	80491857 (2201, 2202) Novel Protein sim. GBank gil1174735 sp P43012 TOP1_HAEIN - DNA TOPOISOMERASE I (OMEGA-PROTEIN) (RELAXING ENZYME) (UNTWISTING ENZYME) (SWIVELASE)		81897259 (2205, 2206) Novel Protein sim. GBank gi[1906596 (U81788) - kinesin-73 [Drosophila melanogaster]	Novel Protein sim. GBank gil2935448 (AF048976) - synaptic ras GTPase-activating protein p135 SynGAP	Constant of the contract of th			Novel Protein sim. GBank gil1173421[sp]P43416[SECY_STRSC - PREPROTEIN TDANSI OCASE SECY SHIRINIT	80064615 (2217, 2218) Novel Protein sim. GBank gil2995310 emb CAA18338  - (AL022258) putative ATP-dependent helicase		Novel Protein sim. GBank gi 2622039 (AE000868) - type I restriction modification system, subunit S Methanobacterium thermoautotrophicum)		82456352 (2225, 2226) Novel Protein sim. GBank gij3218376jembjCAA19628j - (AL023862) putative oxidoreductase (Streptomyces coelicolor)	14988014 (2227, 2228) Novel Protein sim. GBank gil1083428 pir  S54876 - NAD(P)+ transhydrogenase (B-specific) (EC 1.6.1.1)	piecursor - mouse		
1099 39523838 (2197, 2198) Novel Protein sim. GBanl glj3915144 spj033017 TF N1)-METHYL TRANSFERASE METHYLTRANSFERASE METHYLTRANSFERASE METHYLTRANSFERASE	85736571 (2199, 2200) Novel Protein sim. GBank gij3023255jsp Q64420]AC DESATURASE (STEARC ACID DESATURASE) (DI	80491857 (2201, 2202) Ni 91 77 EP	79777614 (2203, 2204)	81897259 (2205, 2206) N	95003115 (2207, 2208) Novel Protein sim. GBan synaptic ras GTPase-ac	R0255121 (2209 2210)	70314110 (2211 2212)	80470019 (2213 2214)			80503554 (2219, 2220)	80071744 (2221, 2222)	95010088 (2223, 2224)		14998014 (2227, 2228)	4476669 (2000 0000)	11/63363 (2229, 2230)	1116 [79841152 (2231, 2232)]
1099	1100	1101	1102	1103	1104	1105	300	1107	1108	1109	1110	1111	1112	1113	1114	4446		1110

18108392, 264488, 263894, 264489, 18108392, 264488, 25278998, 25278998, 25278998, 25278998, 25278998, 25278998, 25278998, 25278998, 25278998, 25278998, 25278998, 25278998, 25278998, 25278999, 264259, 29331822, 56182181, 26331824, 66174117, 29331828, 3659652, 3331826, 264509, 264908, 264508, 264909, 264908, 264907, 264908, 66712502, 264909, 52644045, 56182435, 264510, 264511, 265008, 265007, 264512, 265008, 26509, 264910, 60170831, 264591, 264593, 264593, 264593, 264593, 264601, 264601, 264758, 264769, 264601, 264601, 264764, 265018, 264760, 55811150, 264681, 264762, 264681, 264762, 264681, 264685, 24468, 264884, 264885, 24468, 264687, 264769, 264681, 264682, 264769, 264681, 264682, 264693, 264691, 265021, 265022, 20170615, 264691, 265021, 265022, 264693, 65274620, 33657109, 33657182, 27486261, 27486262, 27486264, 33657192, 27486261, 27486264, 3365739, 27486261, 27486262, 27486264, 33657399, 27486261, 27486264, 33657399, 27486261, 27486264, 33657399, 27486261, 27486264, 33657399, 27486261, 27486264, 33657399, 27486261, 27486264, 33657399, 27486261, 27486264, 33657399, 27486261, 27486264, 33657399, 27486261, 27486264, 33657399, 27486261, 27486264, 33657399, 27486261, 27486264, 33657399, 27486261, 27486264, 33657399, 27486261, 27486261, 27486264, 27486264, 27486261, 27486264, 27486264, 27486261, 27486264, 27486261, 27486261, 27486264, 27486261, 2748621, 27486261, 27486261, 2748621, 27486261, 2748611, 27486	264691	264907	29331827, 265018, 265019, 264681, 265021, 60170615, 18108387	264683, 264632, 18108388	264639, 264563	264693, 27486265	264909	264910	264905, 264909, 264910	264682	264511
helicase	UNCLASSIFIED	UNCLASSIFIED	collagen	UNCLASSIFIED	UNCLASSIFIED	synthase	UNCLASSIFIED	UNCLASSIFIED	eph	UNCLASSIFIED	- polymerase
Contains protein domain (PF00270) - helicase DEAD/DEAH box helicase				Contains protein domain (PF00013) - UNCLASSIFIED KH domain					·		Contains protein domain (PF00476) - polymerase DNA polymerase family A
95305465 (2233, 2234) Novel Protein sim. GBank gij3255965[emb]CAA94089] - (Z70200) U5 snRNP-specific 200kD protein [Homo sapiens] DEAD/DEAH box helicase			79480463 (2239, 2240) Novel Protein sim. GBank gil5420387 emb CAB46679.1  (AJ243459) proteophosphodycan [Lelshmania major]	78471716 (2241, 2242) Novel Protein sim. GBank gil (64450 (U67864) - MEX-3 ICaenorhabditis elegans		78637119 (2245, 2246) Novel Protein sim. GBank gij98800 pir  S17768 - 3- dehydroquinate synthase (EC 4.6.1.3) - Mycobacterium tuberandeis	-		79758914 (2251, 2252) Novel Protein sim. GBank gij138154 spiP03643 VGG_BPPHX - MAJOR SPIKE protein AC PROTEIN (GPG)		Novel Prolein sim. GBank gij5002704 jemb CAB44358.1  - (AJ242630) DNA polymerase I [Methylobacterium sp. DM4]
95305465 (2233, 2234)		79642463 (2237, 2238)		79471716 (2241, 2242)	79456246 (2243, 2244)		79811596 (2247 2248)			11800930 (2253, 2254)	_
2117	1118	1119	1120	1121	1122	1123	1124	1125	1126	127	1128

265011, 264768	264595	35696286, 22278998, 29331828, 264603, 264605, 264559	22278998 264906 265009 264600 264602	264604, 264605, 264760, 32833986, 18108374	265018	264512, 264534	264508, 264600, 264602, 264803, 181083/6	264906, 264908	22278998, 22278999, 35696052, 264907, 265009, 60433356, 264596, 265010, 264448, 264682, 264767, 264689, 265020, 264692, 55811576, 35695855, 264631, 264632, 22279002	264908	264693	264593	265019, 264693	264631	56182575, 264908, 264600, 264632, 87168518	264635, 264936, 264907, 264593, 264908, 264566, 264909	264112	264769, 264689, 35696286, 264760, 264905, 264486, 284559	284557	284591	264591
JNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	INC. ACCIEIED	ONCHASSIFIED	UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	collagen	struct	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	oxidase	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED		kinase	
Contains protein domain (PF00170) - UNCLASSIFIED b2IP transcription factor			5						Contains protein domain (PF00515) - collagen TPR Domain						Contains protein domain (PF00090) - oxidase Thrombospondin type 1 domain						
80422480 (2257, 2258) Novel Protein sim. GBank gil5689485 dbj BAA83026.1  -		80055391 (2261, 2262) Novel Protein sim. GBank	gilago iszelgupAdosogi i pedori i - (Aecon 17) do protein [Thermotoga maritima]	82052248 (2263, 2264) Novel Protein sim. GBank gi 1841552 (U89336) - unknown [Homo sapiens]			80029393 (2269, 2270) Novel Protein sim. GBank gil4539171[emb[CAB39700.1] - (AL049485) conserved hypothetical protein [Streptomyces coelicolor]	79842052 (2271, 2272) Novel Protein sim. GBank gil4982454[gb]AAD36931.1[AE00182 - (AE001823) ATP. dependent protease LA. putative [Thermotoga maritima]	90931557 (2273, 2274) Novel Protein slm. GBank gil4972746 gb AAD3476B.1  - (AF132180) unknown [Drosophila melanogaster]	79841163 (2275, 2276) Novel Protein sim. GBank gij731607[splP38739]YHC8_YEAST - HYPOTHETICAL 63.8 KD PROTEIN IN GUT1-RIM1 INTERGENIC REGION	PRECURSOR 78633561 (2277, 2278) Novel Protein sin. GBank gij3650031 (AC005396) - putative	-			87762158 (2285, 2286) Novel Protein sim. GBank gij3928000jemb CAA05880j - (A_003125) procollagen I N-proteinase [Homo sapiens]					Novel Protein sim. GBan gi[2499003 sp P76422 Ti PHOSPHOMETHYLPYR	11607438 (2297, 2298) Novel Protein sim. GBank gi[2896734[emb]CAA17213.1] - (AL021897) hypothetical protein Rv1097c [Mycobacterium tuberculosis]
10422480 (2257, 2258)	10400 00EG (100EG)	30055391 (2261, 2262)		32062248 (2263, 2264)	17290437 (2265 2268)	80235376 (2267, 2268)	80029393 (2269, 2270)	79842052 (2271, 2272)	90931557 (2273, 2274)	79841163 (2275, 2276)	79633561 (2277, 2278)	39480358 (2279, 2280)	79638019 (2281, 2282)	19635848 (2283, 2284)	87762158 (2285, 2286)	80088988 (2287, 2288)	14610262 (2289, 2290)		80071761 (2293, 2294)		
1129 8	_	1131	_	1132 8	-	1134		1136	1137	1138	1139	1140	1141	1142	1143	1144	1145	1146	1147	1148	1148

ATPase_associated 264488, 35696286, 264907, 264908, 264909, 264909, 264910, 284593, 264596, 264758, 264764, 264768, 284768, 264693, 264628, 60431850.	264595	264488, 22278998, 264905, 264629, 264486	264910, 264555, 264557	265008	254603	200403	264906, 264907, 264758, 264768, 264769. 264689, 264638, 264566	26490B 264762 264687 264769 264689.	18108374, 35695855	264687	2931822, 2931824, 66/14117, 2951626, 60433356, 265018, 265019, 83373044, 18108385, 22279000, 22279002, 264563	264369	264693	29331827, 264906	18108398, 29331827, 29331829, 29146499, 29146499, 18108354, 21906768, 29148627, 21906769, 264693, 18108382, 18108385	264602, 264605, 264769, 18108370, 18108374, 264565	264488, 35696286, 22278999, 264259, 66714117, 60432289, 35696052, 264905, 56182435, 265006, 60433438, 264759, 21906754, 33109954, 265017, 265019,	26448, 264288, 264766, 264685, 35696423, 35695855, 264558, 18108385, 60432113
ATPase_associated	polymerase	glycoprotein		UNCLASSIFIED	UNCLASSIFIED	lranspor	UNCLASSIFIED						UNCLASSIFIED	UNCLASSIFIED	transport	UNCLASSIFIED		
						Contains protein domain (Pr00122) - Iransport E1-E2 ATPase									,			
1150   81325074 (2299, 2300) Novel Protein sim. GBank gij2895095 (AF011337) - putative E1-E2 ATPase (Mus musculus)	80070874 (2301, 2302) Novel Protein sim. GBank giļ4324655[gb[AAD16978] - (AF108191) DNA polymerase lil alpha subunit	Streptomyces coelicolor    Streptomyces coelicolor    Novel Protein sim. GBank gij3874275 emb CAB07311.1 -   (292825) predicted using Genefinder: Similarity to Yeast low afinity glucose transporter HT74 (PS:32467); cDNA EST EMBL:C12555 comes from this gene; cDNA EST yk404c10.5	comes irom (ni	Novel Protein sim. GBank gi[4240315]dbj BAA74936.1  - (AB020720) KIAA0913 protein [Homo sapiens]		57147843 (2311, 2312) Novel Protein sim. GBank gil586655 sp P37617 ATZN_ECOLI - ZINC- TRANSPORTING ATPASE (ZN(II)-TRANSLOCATING P-	95287711 (2313, 2314) Novel Protein sim. GBank gil418480 sp P32139 YIHR_ECOLI - HYPOTHETICAL 34.0	KD PROTEIN IN GLNA-RBN INTERGENIC REGION	82454917 (2315, 2316) Novel Protein sim. GBank gi 2496481 sp 050724 Y09S_MYCTU - HYPOTHETICAL  87.3 KD PROTEIN CY78.27C		91229893 (2319, 2320) Novel Protein sim. GBank gill136406 dbj BAA11490  - (D79995) similar to pig tubulin-tyrosine ligase. [Homo	Cipidac	79635357 (2323, 2324) Novel Protein sim. GBank gi[2443342 dbj BAA22380  - 79635357 (2323, 2324) Novel Protein sim. GBank gi[2443342 dbj BAA22380  - 70635357 (2323, 2324) Novel Protein sim. GBank gi[2443342 dbj BAA22380]	79563186 (2325, 2326) Novel Protein sim. GBank Initegrary Flipeting 001376 11DDPYS - dibydropyrimidinase	79650829 (2327, 2328) Novel Protein sim. GBank gij5052554[gb]AAD38607.1 AF14563 - (AF145632)	BCDNA, GHU6032 [Urosopnila melandgaster]	88096456 (2331, 2332) Novel Prolein sim. GBank gil4589476 dbj BAA76766.1  - (AB023139) KIAA0922 protein [Homo sapiens]	
1325074 (2299, 2300) N	0070874 (2301, 2302) N.	0235547 (2303, 2304) N (7	0000 3000 60750000	+=-	79411098 (2309, 2310)	57147843 (2311, 2312) N	95287711 (2313, 2314)		82454917 (2315, 2316)	10100 710011	91229893 (2319, 2320)	1447449 (9994 9999)	1=	79563186 (2325, 2326)	79650829 (2327, 2328)	80491888 (2329, 2330)		
1150 8	1151 80	1152 8		1154	-	1156	1157		1158		1160		1162	1163	1164	1165	1168	· 

264488	264259, 29331827, 56182435, 60433438, 265019, 33657023, 35695855, 264566	264681	264602	284556	55810764, 35696052, 264634, 264486	264508, 264906, 264907, 264909, 264591, 284632, 264638, 264839	264600	264601	264509, 264905, 264593, 264602, 264605		65274572, 18108398, 22278998, 22278999, 29331826, 284508, 284908, 284808, 284808, 284808, 284808, 284828, 33109954, 284769, 21906765, 21906768, 55811957, 33657023, 264629, 55811576, 35694421, 284856, 56182323, 60432113, 22279000, 22279002	264603	264638	265017	265007, 265009, 264564, 264909, 264693	264689, 35696423, 264638, 18108385	264802	263967
	UNCLASSIFIED	UNCLASSIFIED	П		UNCLASSIFIED		se		synthase	reductase	ATPase_associated	transport	synthase	UNCLASSIFIED			UNCLASSIFIED	glycoprotein
	**							Contains protein domain (PF00205) - carboxylase Thiamine pyrophosphate enzymes	Contains protein domain (PF00586) - synthase AIR synthase related protein	Contains protein domain (PF00106) - reductase short chain dehydrogenase	Contains protein domain (PF00122) - ATPase_associated E1-E2 ATPase	Contains protein domain (PF00496) - transport Bacterial extracellular solute-binding proteins, family 5	Contains protein domain (PF01220) - synthase Dehydroquinase class II					
79963862 (2333, 2334) Novel Protein sim. GBank gi 2580433 db  BAA23138  -  (D76414) ppGpp hydrolase  Staphylococcus aureus)				20434582 (2341, 2342) Novel Protein sim. GBank gil2772914 (AF029249) - precollagen D (Mytllus edulis)	79610113 (2343, 2344) Novel Protein sim. GBank gil4757846[refINP_004317.1[pBCL9 - 8-cell CLUlymphoma	Novel Protein sim. GBank gi[2564053 dbj BAA22946  - ,	20293077 (2347, 2348) Novel Protein sim. GBank gil2911027[emb CA417520] - IAI 021958) mmsA IMvcobacterium tuberculosis	20711847 (2349, 2350) Novel Protein sim. GBank gil118333 sp P23234 DCIP_ENTCL - INDOLE-3- PYRUVATE DECARBOXYLASE (INDOLEPYRUVATE DECARBOXYLASE)	80252645 (2351, 2352) Novel Protein sim. GBank gi[1144520 (U34956) - phosphoribosylformyglycinamidine synthase [Mycobacterium tuberculosis]	80064647 (2353, 2354) Novel Protein sim. GBank gij 19791 spjP28643jFABG_CUPLA - 3-OXOACYL-{ACYL- CARRIER PROTEIN] REDUCTASE PRECURSOR (3- KETOACYL-ACYL CARRIER PROTEIN REDUCTASE)	94128641 (2355, 2356) Novel Protein sim. GBank gi 5031697 ref NP_005594.1 pFIC1 - familial intrahepatic cholestasis 1. (progressive, Byler disease and benign recurrent)	80055575 (2357, 2358) Novel Protein sim. GBank gil2860090 emb CAA17988.1 - (AL022121) dppA [Mycobacterium tuberculosis]	11794448 (2359, 2360) Novel Protein slm. GBank gij2558614 emb CA004787 - (AJ001493) dehydroquinate dehydralase (Streptomyces	_	81494284 (2363, 2364) Novel Protein sim. GBank gij5420387jembjCAB46679.1  - (A_243458) proteophosohodycan IL eishmanla maiori		52559933 (2367, 2368) Novel Protein sim. GBank gil4091877 (AF061331) - alpha qalactosidase precursor [Saccharopolyspora erythraea]	79491185 (2369, 2370) Novel Protein sim. GBank gij2129478[pirj[551939 - chitinase (EC 3.2.1.14) precursor - beet
79963862 (2333, 2334) N	88094678 (2335, 2336)	11805403 (2337 2338)	21632244 (2339, 2340)		79610113 (2343, 2344) N	80235713 (2345, 2346) N	20293077 (2347, 2348) h	20711847 (2349, 2350) h	80252845 (2351, 2352) P	80064647 (2353, 2354)    	94128641 (2355, 2356)	80055575 (2357, 2358)		17946362 (2361, 2362)	81494284 (2383, 2384)	78574044 (2365, 2366)	52559933 (2367, 2368)	_
1167 7	1168	1169 1	_	_	1172 7	1173	1174	1175	1178	1177	1178	1179	1180	1181	1182	1183	184	1185

1186	20224012 (2371 2372)		1	INCI ASSIFIED	264559
	79248834 (2373, 2374)			UNCLASSIFIED	29331825, 265017, 18108351
1188	79831387 (2375, 2376)	79831387 (2375, 2376) Novel Protein sim. GBank gl 2996039 (AF054525) - hypothetical protein [Synechococcus PCC7002]		UNCLASSIFIED	264905, 264908
1189	79609367 (2377, 2378)				264692
1190	78930589 (2379, 2380)			UNCLASSIFIED	265018
1191	80310105 (2381, 2382)			UNCLASSIFIED	264600, 264605, 264764, 35695855, 264638, 264486
1192	13521641 (2383, 2384)		Contains protein domain (PF01581) - FMRFamide related peptide family		264636
193	11103584 (2385, 2388)		Γ	UNCLASSIFIED	263978
1194	78893947 (2387, 2388)	78893947 (2387, 2388) Novel Protein sim. GBank gil854065 emb CAA58337  - (X83413) U88 [Human herpesvirus 6]		UNCLASSIFIED	265007, 265008
1195	20445442 (2389, 2390)			UNCLASSIFIED	264605
1196	13000688 (2391, 2392)				264689
197	11392317 (2393, 2394)	11392317 (2393, 2394) Novel Protein sim. GBank	Contains protein domain (PF00571) - dehydrogenase	dehydrogenase	264594
		gij2497360jsp Q50715jiMDH_MYCTU - INOSINE-5: MONOPHOSPHATE DEHYDROGENASE (IMP DEHYDROGENASE) (IMPDH) (IMPD)	CBS domain		
1198	95290101 (2395, 2396)				264603
199	_	81882011 (2397, 2398) Novel Protein sim. GBank			264259, 264757, 33109954, 21906768
		gij 709525 sp P54673 P3K1_DICDI - PHOSPHATIDYLINOSITOL 3-KINASE 1 (P13-KINASE) (PTDINS-3-KINASE) (P13K)			
1200 1200	_			UNCLASSIFIED	264910
201		80503751 (2401, 2402) Novel Protein sim. GBank gi 2499877 sp P70645 BLMH_RAT - BLEOMYCIN HYDROLASE (BLM HYDROLASE) (BMH)		cathepsin	264766, 264769
1202		80082633 (2403, 2404) Novel Protein sim. GBank gi[606342 (U18997) - ORF_0622; reading frame open far upstream of start; possible frameshift, linking to previous ORF [Escherichia coli]		ribosomalprot	264600, 264558
1203	82125373 (2405, 2406)			UNCLASSIFIED	264768, 264769, 35695917, 264910, 264760, 264909, 264909, 264569, 264909, 264568
1204		80503916 (2407, 2408) Novel Protein sim. GBank gi 2500728 sp 059912 SECY_STRGB - PREPROTEIN TRANSLOCASE SECY SUBUNIT			264905, 264769, 264636
1205				UNCLASSIFIED	264566
1208	_		,	UNCLASSIFIED	264556, 264557, 264558
<b>1</b> 20					29331824, 264909, 265021, 18108370
88 28		87755217 (2415, 2416) Novel Protein sim. GBank gi 2645560 (AF027954) - Bct-2- related ovarian killer protein [Rattus norvegicus]	Contains protein domain (PF00452) - Apoptosis regulator proteins, Bcl-2 family	apoptosis	29331824, 29331825, 29331827, 265007, 264764, 264683, 264769, 264688, 264689

264687, 264688	264907, 264693	264591, 284592, 264595	264689, 263967	264909	264908	265009, 264601, 264602, 264603, 3365/109	264604	264905, 264907, 264908, 264591, 264766, 264691, 264693, 264629, 264630, 264636, 264564	264601	56181686, 29331822, 60432289, 264601, 264692, 264629	52644507, 264905, 264909, 265008, 265019, 265020, 52644150, 33657023, 264693, 33657182, 35695763, 264634, 22279000, 22279002, 264482	264768, 265020, 264906		264593	264593, 264600, 264601, 264603, 264605, 264768, 18108376, 264635, 18108387
isomerase	UNCLASSIFIED	transferase		kinase	UNCLASSIFIED	UNCLASSIFIED	transferase	UNCLASSIFIED	dehydrogenase		oncogene	UNCLASSIFIED	- phosphatase		- kinase
Contains protein domain (PF00259) - isomerase Xytose isomerase			Contains protein domain (PF01412) - Putative GTP-ase activating protein for Arf										Contains protein domain (PF00486) - phosphatase Transcriptional regulatory protein, C terminal		Contains protein domain (PF01472) - kinase PUA domain
sím. GBank IP44398JXYLA_HAEIN - XYLOSE	ISOMEKASE	30420304 (2415, 2420) 94665655 (2421, 2422) Novel Protein sim. GBank gil421095[pir][530688 -	gij3880625jembjCAB07858j - Genefinder; similar to RNA KM, RBD, or RNP domain); cDNA s from this gene; cDNA EST m this gene; cDNA EST m this gene; cDNA EST	78859633 (2425, 2426) Novel Protein sim. GBank gi[226292 prt] 1505375A - vir gene [Bordetella pertussis]	10144306 (2427, 2428) Novel Protein sim. GBank gij5726285[gb]AAD48396.1]AF12616 - (AF126162) HERV-H LTR associating protein 2 [Homo sapiens]	80050108 (2429, 2430) Novel Protein sim. GBank gil2326739[emb]CAB10953  - IZ28258) recN (Mycobacterium tuberculosis)	20438324 (2431, 2432) Novel Protein sim. GBank gij417328jspjP33038jMURA_ENTCL - UDP-N- ACETYLGUCOSAMINE 1- CARBOXYVINYLTRANSFERASE (ENOYLPYRUVATE TRANSFERASE) (UDP-N-ACETYLGUCOSAMINE FROID PYRUXY TRANSFERASE) (FPT)		11093680 (2435, 2436) Novel Protein sim. GBank gi 1805460 db  BAA09022 - (D50453) homologue of succinate semialdehyde dehydrocenase GabD of E. coli [Bacillus subfilis]		91241524 (2439, 2440) Novel Protein sim. GBank gil4240315 dbj BAA74936.1 - (AB020720) KIAA0913 protein [Homo sapiens]	83045055 (2441, 2442) Novel Protein sim. GBank gij2143886 pir  152523 - nucleoporin p82 homolog - rat (fragment)	20711865 (2443, 2444) Novel Protein sim. GBank gij730805[spiP39663 SPHR_SYNP7 - ALKALINE PHOSPHATASE SYNTHESIS TRANSCRIPTIONAL REGIJJ ATORY PROTEIN SPHR	-	80432845 (2447, 2448) Novel Protein sim. GBank gij1172627 sp P46546 PROB_CORGL - GLUTAMATE 5- KINASE (GAMMA-GLUTAMYL KINASE) (GK)
79185742 (2417, 2418	C6420 0440 4400	94665655 (2421, 2422	79167929 (2423, 242 <sup>,</sup>	79859633 (2425, 242)	_			95011344 (2433, 2434)	+	91216252 (2437, 2438)		_	7	11615647 (2445, 2446)	
1209	355	1211	1212	1213	1214	1215	1216	1217	1218	1219	1220	1221	1222	1233	1224

1225	1225 80434427 (2449, 2450)				264768
1228	80237518 (2451, 2452)	1226   80237518 (2451, 2452)   Novel Protein sim. GBank gil2105050 emb CAB08836  - (295436) hypothetical protein Rv3644c   Mycobacterium		polymerase	264905, 264512, 264689
1227	79422138 (2453, 2454)	79422138 (2453, 2454) Novel Protein sim. GBank 911706768 sp P98133 FBN1_BOVIN - FIBRILLIN 1 PRECURSOR (MP340)		UNCLASSIFIED	264908, 264637, 264639
1228	79209027 (2455, 2456)	79209027 (2455, 2456) Novel Protein sim. GBank gil1653901 dbjlBAA188111- (D90917) acrifavine resistance protein (Synechocystis sp.)	Contains protein domain (PF00873) - Acr8/AcrD/AcrF family		264605, 264634
1229	94329135 (2457, 2458)			UNCLASSIFIED	87168474, 265011, 87168559, 264681, 264689, 264693, 65274620, 18108374
1230		80049357 (2459, 2460) Novel Protein sim. GBank gil116230 sp P28598 CH60_BACSU - 60 KD CHAPERONIN TCP-1/cpn60 chaperonin family (PROTEIN CPN80) (GROEL PROTEIN)	Contains protein domain (PF00118) - eph TCP-1/cpn60 chaperonin family	hdə	264909, 264605, 18108388
1231	79843141 (2461, 2462)			UNCLASSIFIED	264908
1232		79853104 (2463, 2464) Novel Protein sim. GBank gi[1215733 (U48718) - OphC [Agrobacterium tumefaciens]		transport	264909
1233	80255179 (2465, 2466)	80255179 (2465, 2466) Novel Protein sim. GBank		UNCLASSIFIED	265017, 264564
		gi[116298[sp P20730]CHHC_BOMMO - CHORION CLASS HIGH-CYSTEINE HCB PROTEIN 13 PRECURSOR (HC- 18.13)			,
1234	79242158 (2467, 2468)	Protein sim. GBank	Contains protein domain (PF00125) - histone	histone	265008, 265010, 18108381
1226	70014473 (7460 2470)	gilizabi ilspirauzoninza maize - nis i ONE HZA	Core nistone HZAVHZB/H3/F4		001001 001260
1220	_				204034, 204/02
2	0192/14/ (24/1, 24/2)			UNCLASSIFIED	265018, 55811150, 264565, 264757
1237		833/1782 (2473, 2474) Novel Protein sim. GBank gij3875133 emb CAA94750j - (270750) similar to actin binding domain; cDNA EST EMBL.T00093 comes from this gene; cDNA EST EMBL.D34443 comes from this gene; cDNA EST EMBL.D37508 comes from this gene; cDNA EST EMBL.D37508 comes from this gene; cDNA EST EMBL.D84247 comes from this gene; cDNA EST			264758, 284561, 264766, 264687, 18108372, 264555, 264559
1238	87411577 (2475, 2476)	87411577 (2475, 2476) Novel Protein sim. GBank gij3885470 (AF061443) - G protein-coupled receptor LGR4 [Rattus norvegicus]	Contains protein domain (PF00560) - glycoprotein Leuclne Rich Repeat	glycoprotein	264259, 29331822, 29331824, 35696052, 264508, 264906, 52644045, 52646317, 264288, 264769, 264693, 264632, 264634, 264558, 87168518, 264563
1239		82197449 (2477, 2478) Novel Protein sim. GBank gil4007990 gblAAC95339  -  (AF084383) DOK protein [Mus musculus]		oncogene	284509, 264511, 264759, 264760, 264764. 264557
1240					264769
1241			Contains protein domain (PF00120) - UNCLASSIFIED Glutamine synthetase	UNCLASSIFIED	264601, 264604, 264638
1242	79775890 (2483, 2484)				264906, 264907, 264908, 264634

				ſ	
1243	797 79458 (2485, 2486)	1243   /97/9458 (2485, 2486) Novel Protein sim. GBank gij355567 jembjCAA19971  - (AL031124) branched-chain amino acid aminotransferase [Streptomyces coelicotor]		UNCLASSIFIED	181083/4, 35695817, 35695855, 265009, 264508, 264909
		10284821 (2487, 2488) Novel Protein sim. GBank gil2970646 (AF051945) - Xin [Mus musculus]		UNCLASSIFIED	264691
1245		Novel Protein sim. GBank gil4586338 dbj BAA76357.1  - (AB016787) cytochrome o ubiquinol oxidase B [Pseudomonas putida]	Contains protein domain (PF00115) - oxidase Cytochrome C and Quinol oxidase polypeptide I		264768
1248		80059321 (2491, 2492) Novel Protein sim. GBank gij3581849 emb CAA20805] - (AL031541) putative phenylalany-IRNA synthetase beta chain [Streptomyces coelicotor]		UNCLASSIFIED	264604, 264636, 264557, 264564
1247	80064831 (2493, 2494)	80064831 (2493, 2494) Novel Protein sim. GBank gilz621684 (AE000842) - adhesion protein (Methanobacterium thermoautotrophicum)			264758, 264605, 264639
1248	88070353 (2495, 2496)	88070353 (2495, 2496) Novel Protein sim. GBank gij1352403jspp09467 F16P_HUMAN - FRUCTOSE-1,6- BISPHOSPHATASE (D-FRUCTOSE-1,8-BISPHOSPHATE 1-PHOSPHOHYDROLASE) (FBPASE)	Contains protein domain (PF00316) - UNCLASSIFIED Fructose-1-6-bisphosphatase	UNCLASSIFIED	18108392, 264259, 29331826, 264106, 264508, 264508, 264907, 264828, 265009, 60433356, 264757, 264758, 21906754, 265010, 265011, 265018, 265019, 264750, 18108351, 18108354, 265021, 18108376, 18108377, 264630, 18108385
1249	80056657 (2497, 2498)	80056657 (2497, 2498) Novel Protein sim. GBank gi[2791407 emb CAA16001  - (AL021184) hypothetical protein Rv1473 [Mycobacterium [tuberculosis]		transport	264908, 265010, 264600, 264603, 264691, 18108376
1250	12694385 (2499, 2500)	1250 12694385 (2499, 2500) Novel Protein sim. GBank gi[112785 sp P05100 3MG1_ECOLI - DNA-3- METHYLADENINE GLYCOSIDASE I (3-METHYLADENINE- DNA GLYCOSYLASE I, CONSTITUTIVE) (TAG I)		UNCLASSIFIED	264689
1251	_				264909
1252	_			ED	264683, 263976
1253		80050121 (2505, 2506) Novel Protein sim. GBank gil5670176[gb]AAD46616.1JAF16131 - (AF161317) NRAMP manganese transport protein MnLA (Salmonella (typhimurium)	-1		264600, 264603, 18108376
1254		87716767 (2507, 2508) Novel Protein sim. GBank gi 103160 pir  S22126 - finger protein unkempt - fruit fly (Drosophila melanogaster)		UNCLASSIFIED	35696286, 264310, 264764, 264688, 21906767, 55811957, 264692, 264556, 264639
1255	-			•	264636
1256		87889508 (2511, 2512) Novel Protein sim. GBank gi 2995353 emb CAA04608.1  -  (AJ001206) pep2 [Streptomyces coelicolor]		UNCLASSIFIED	60432289, 264600, 264605, 264764, 264687, 264769, 264689, 27486265, 18108374, 18108376
1257		80201435 (2513, 2514) Novel Protein sim. GBank gij3193306 (AF069300) - contains similarity to Arabidopsis membrane-associated salt inducible-like protein (GB:AL021637) [Arabidopsis thaliana]	. =	UNCLASSIFIED	264094, 265019
1258				UNCLASSIFIED	264602, 263978
1259	80186012 (2517, 2518)			UNCLASSIFIED	264906, 264448, 264908
1280				UNCLASSIFIED	264634, 264639

29331824, 264906, 264909, 264768, 264769, 264689, 264693, 264639, 18108384, 264563	264634	264602			264563	264766, 264636, 264638, 264567	264259, 21906754, 264369		18108398, 22278995, 56994075, 60424269.	29331827, 264109, 264512, 265007, 265008.	265009, 264595, 33109954, 33657084.	87168559, 264600, 265018, 265019, 264369,	264688, 21906767, 265020, 52644150.	264691, 33657023, 33657349, 18108374.	264556, 18108385, 60432113, 22279002,	204400	264508, 264805, 264806, 264667, 264655		265010, 264601		ATPase_associated 35696052, 264908	264686, 264689	264600 28483Q	60000, 600000	264905, 264908, 264909, 264769	264602	70000		264369		264556	264603		264605	00000	0/8787
	UNCLASSIFIED		Profession of the Profession o	Don No.	UNCLASSIFIED	UNCLASSIFIED	fransport		UNCLASSIFIED					٠			UNCLASSIFIED		dehydrogenase		ATPase_associated		000000000000000000000000000000000000000	UNCLASSIFIED	INCI ASSIFIED	4-5	ebu_		UNCLASSIFIED	1-	UNCLASSIFIED	UNCLASSIFIED		synthase		struct
Contains protein domain (PF01699) - cadherin Sodium/calcium exchanger protein							PENNIS Services domain (PENN153) - Iransport	Mitochondrial carrier proteins	Contains profein domain (PF00583) - UNCLASSIFIED	Acetyltransferase (GNAT) family			,						Contains protein domain (PF01574) - dehydrogenase	IMP dehydrogenase / GMP						6616	Contains protein domain (PF00183) - epn	Hsp90 protein								
13039.1 -	13504589 (2523, 2524) Novel Prolein sim. GBank gil95100 pri S21334	hypothetical protein 4 - Agrobacterium tumefaciens		80083386 (2527, 2528) Novel Protein sim. GBank gij3550958 (AF004840) - CDO IIRatius norveolousì				80558918 (2533, 2534) Novel Protein sim. GBank gij 108500Zipirijsəsəbəb - mitochondrial carner protein DIF-1 homolog -	Caendragouis elegaiis	88178473 (2535, 2536) Novel Protein sim. Gbank gipoco443jeniojovov. (1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1								(AL031107) hypothetical protein SC3A7.1UC (Streptomyces	Cochical Service Cochical Coch	gilgasidasin sim. Obdin. gilgasidasidasidasidasidasidasidasidasidasid				_	tuberculosis		20730763 (2549, 2550) Novel Protein sim. GBank	gij123726ispiP10413jHTPG_ECOLI - HEAT SHOCK PROTEIN HTPG (HIGH TEMPERATURE PROTEIN G)	(HEAT SHOCK PROTEIN COS.3)	21148644 (2551, 2552) Novel Protein Sim. Gbank gijd 1294 (9jbi jjbu 1959 - Ichiinase (FC 3.2.1.14) preduksor - heet	-		11088383 (2555, 2550) Novel Fridein sim. Obain gill/134731919194555[YAA]_HAEIN - HYPOTHETICAL	21658756 (2557, 2558) Novel Protein sim. GBank qi 1929513 (U64318) - ATP	synthase subunit beta (Moorella Ihermoacelica)	79310959 (2559, 2560) Novel Protein sim. GBank gil4938504(emb CAB43862.1     (AL078465) putative protein [Arabidopsis thaliana]
7412802 (2521, 2522)	3504589 (2523, 2524)	-	20710997 (2525, 2526)	10083396 (2527, 2528)	02520 72570 2520	80233378 (2328, 2330)	79914604 (2531, 2532)	30558918 (2533, 2534)		88178473 (2535, 2536)							79821946 (2537, 2538)		100000 00000000000000000000000000000000	60031420 (2539, 2540)	C130 11301 0010100	18040488 (2341, 4342)	79462878 (2543, 2544)	80220315 (2545, 2546)		95010802 (2547, 2548)	20730763 (2549, 2550			21148644 (2551, 2552	20438105 (2553 2554)			_	_	
1261 8	1262 1	_		1264 8	_	607	_	1267 8	_	1268						_	1269			0/21	_	7	1272			1				1276	1777		12/8	1279		1280

1281	94323988 (2561, 2562)		Contains protein domain (PF00047) - UNCLASSIFIED Immunoolobulin domain	UNCLASSIFIED	29331825, 29331828, 264766, 83373044
	87537695 (2563, 2564)	87537695 (2563, 2564) Novel Protein sim. GBank gil3328190 (AF074266) - proto- oncogene AF4 (Mus musculus)		UNCLASSIFIED	265008
	20466305 (2565, 2566)	20466305 (2565, 2566) Novel Protein sim. GBank gil3261721 emb CAB07057  - (Z92770) hypothetical protein Rv0153c {Mycobacterium (tuberculosis}		UNCLASSIFIED	264605
	20636325 (2567, 2568)	20636325 (2567, 2568) Novel Protein sim. GBank gij3929022 (AF057696) - LspB [Haemophilus ducreyi]			264604
1285	80427330 (2569, 2570)		Contains protein domain (PF00183) - eph Hsp90 protein	udə	264766, 264689, 263967
1286	20465254 (2571, 2572)	20465254 (2571, 2572) Novel Protein sim. GBank gi 2078004 emb CAB08451  -   (295207) gorA [Mycobacterium tubercutosis]		reductase	264605, 264639
1287				UNCLASSIFIED	265011, 264602, 264766, 284687, 264769, 264689, 18108370, 264638, 18108385, 264563
1288		95338101 (2575, 2576) Novel Protein slm. GBank gi 5353510 gb AAD42161.1 AF08891 - (AF088916) emilin precursor [Homo sapiens]	Contains protein domain (PF00386) - collagen C1q domain	collagen	35696052, 264107, 264508, 264509, 264905, 264906, 264907, 264908, 284909, 264510, 264511, 265007, 264512, 264910, 265009,
	·				3355/402, 264595, 264756, 265011, 265019, 264760, 18108351, 264681, 264764, 264288, 264685, 264766, 264687, 264768, 264769, 265020, 265021, 264534, 264692, 18108370, 264588, 264558, 18108385, 264564, 264568, 264567, 264486, 18108381, 264564,
1289				UNCLASSIFIED	264637
1290		19526027 (2579, 2580) Novel Protein sim. GBank gif1169995 sp P46023 GPCR_LYMST - G-PROTEIN COUPLED RECEPTOR GRL101 PRECURSOR	Contains protein domain (PF00001) - tm7 7 transmembrane receptor (rhodopsin family)	tm7	264563
1291	80470266 (2581, 2582)	80470266 (2581, 2582) Novel Protein sim. GBank gi 2072674 emb CAB08305  - (295120) mIE [Mycobacterium tuberculosis]	Contains protein domain (PF00271) - helicase Helicases conserved C-terminal domain	helicase	265007, 265008, 264769
1292	94723316 (2583, 2584)	1292   94723316 (2583, 2584) Novel Protein sim. GBank gil 1835755 (U86338) - zinc finger protein Png-1 [Mus musculus]	gil 1835755 (U86338) - zinc finger  Contains protein domain (PF01530) - Iranscriptfactor Zinc finger, C2HC type	transcriptfactor	264092, 264259, 29331822, 29331824, 264508, 264908, 264909, 264512, 265008, 265009, 284591, 265019, 264369, 264288, 264686, 264768, 264693, 18108374, 264632, 56182323, 264639, 83373044, 22278002, 264482, 264563
1293				UNCLASSIFIED	265006, 55812038, 264369, 264556
1294.	82125908 (2587, 2588)	82125908 (2587, 2588) Novel Protein sim. GBank gi[2129173]pin[JF64453 - oxaloacetate decarboxylase (EC 4.1.1.3) alpha subunit - Methanococcus jannaschii		biotindep	264602, 264605, 264760, 18108351, 264689. 33657023, 264559
1285	11686851 (2589, 2590	1285   11686851 (2589, 2590) Novel Protein sim. GBank gij5441779 emb CAB46803.1		dehydrogenase	264689

1687904	11687904 (2591, 2592)	592)			UNCLASSIFIED	264591, 264639
78639300 (2593, 2594) Novel Protein sim. GBank gil4982191]gbJAAD36686. directed DNA polymerase	(4) (9) (9)	0	1/AE00180 - (AE001805) DNA- I [Thermotoga maritima]	Contains protein domain (PF01367) - polymerase 5-3' exonuclease	polymerase	264693
94239506 (2595, 2596) Novel Protein sim. GBank gene product (Caenorhabo	N (96	ᇣᅋ	Novel Protein sim. GBank gi[1943770 (U97191) - F53F10.1 gene product (Caenorhabditis elegans)		struct	18108348, 265017
80255378 (2597, 2598)	8	l i				264488, 264906, 264909, 22279002, 284586
80064867 (2599, 2600) Novel Protein sim. GBank   R31665_2 [Homo saplens	00	유원	gi 3445181 (AC005498) -	Contains protein domain (PF01352) - transcriptfactor KRAB box	vanscriptfactor	264605
7939614 (2601, 2602)	[20	اگ کے ق	Novel Protein sim. GBank gil4062973 dbi BAA36204.1 - (AB017138) alpha subunit of malonate decarboxylase [Pseudomonas putida]		UNCLASSIFIED	264906
95416198 (2603, 2604)	3	1				85658542, 265020
9684121 (2605, 2606)	<u></u>	1				264908
79377196 (2607, 2608)	<u>e</u>				UNCLASSIFIED	264508
19905899 (2609, 2610)	<u>6</u>	L				264566
13069230 (2611, 2612) Novel Protein sim. GBank (292669) hypothetical pro [luberculosis]	12)	ZÜE	Novel Protein sim. GBank gij3242273jemb CAB07017  - (292669) hypothetical protein Rv0236c [Mycobactertum tuberculosis]		UNCLASSIFIED	264636
82201029 (2613, 2614)	4	L			UNCLASSIFIED	264907, 264592, 264764
1,426814 (2615, 2616	16	ž g	21426814 (2615, 2616) Novel Protein sim. GBank gil 1602421 (U59433) - 3-ketoacyl acyl carrier protein reductase (Bacillus subtilis)	gij1502421 (U59433) - 3-ketoacyl Contains protein domain (PF00516) - reductase sse [Bacillus subtilis] Envelope glycoprotein GP120	reductase	264555
9263011 (2617, 2618	18	e K			transport	264906, 18108354
.0466319 (2619, 2620	R	ž <u>₹</u> 8	20466319 (2619, 2620) Novel Protein sim. GBank gij5459220[emb CAB48893.1] - (AL096837) putative iron-sulfur protein (Streptomyces coelicolor A3(2))		UNCLASSIFIED	264605
87613142 (2621, 2622) ',	22					35696286, 29331827, 264908, 265008, 264764, 264766, 264688, 21906767, 21906769, 35695917, 284691, 264693
18061720 (2623, 2624)	12	ž & ß	88061720 (2623, 2624) Novel Protein sim. GBank gil4455118 gb AAD21084  - (AF125158) zinc finger DNA binding protein 99 [Homo sapiens]	Contains protein domain (PF00096) - dna_ma_bind Zinc finger, C2H2 type	dna_ma_bind	22278995, 22278998, 22278999, 264905, 264908, 265011, 265017, 265019, 264687, 21906768, 265020, 265021, 33657023, 22279007, 244564
11225458 (2625 2626	18	Ž	91225458 (2625, 2625) Novel Protein sim GRank	Contains profess domain (PE00886) - phosomatorot	ribocomalorot	22278996 22278999 264259 20281099
	3	<u></u>	gi[4929733]gb[AAD34127.1 AF15189 - (AF151890) CGI-132 Ribosomal protein S16	Ribosomal protein S16		29146498, 264508, 264908, 66712502,
		<u>ā.</u>	protein [Homo sapiens]			60433356, 60433438, 265011, 265017, 284683 264288 21906765 21906767
						29148627, 21906768, 35695917, 265021,
						33657023, 33657109, 18108370, 18108377,
						35695855, 60432113, 22279000, 264563, 18108390
56926053 (2627, 2628)	28	L	56926053 (2627, 2628)			264693
84357192 (2629, 2630	8	ZE	Novel Protein sim. GBank gl 2589223 (AF026565) - ring linger protein [Mus musculus]	Contains protein domain (PF00097) - interleukinrecept Zinc finger, C3HC4 type (RING Innger)	· interteukinrecept	264691
	١					

56182575, 56181686, 20281171, 29331822, 29331824, 60424269, 29331825, 35696052, 5264045, 264591, 6043229, 265018, 265019, 55811150, 56181562, 21906765, 21906768, 35695917, 60170615, 33657023, 65274620, 33657109, 35695763, 35695655, 18108387, 87168518, 60432113, 22279002, 264564	264093, 264906, 264909, 264369, 264684	18108392, 18108348, 265011, 265017, 18108359, 18108362, 56182323, 18108385, 22279000	264908, 264909, 265006, 265008, 294592, 265019, 264768, 56181562, 18108368, 264628, 264629, 18108377, 264638	35696286, 55812038, 265018, 21906768, 265020, 263978, 22279002	60432049, 28331828, 264907, 264908, 264909, 264909, 264910, 55812038, 264601, 264762, 264764, 264768, 264769, 264628, 18108374, 264634, 264635, 18108385	29331828, 264908, 265020, 33657023, 264693, 264404	65274572, 56994075, 264259, 29331822, 29331827, 264104, 56182435, 87168474, 18108351, 264288, 21906766, 21906767, 35695917, 265020, 264693, 65274791, 56182323, 18108387	22278996, 22278998, 29331828, 264905, 264907, 29331830, 264908, 264907, 29331830, 264908, 264510, 265008, 264595, 264759, 21906754, 265018, 264288, 264768, 264769, 21906768, 265022, 18108376, 264631, 264632, 264565, 264566	35636286, 264905, 264906, 264907, 264908. 264909, 264910, 264593, 33657402, 264758, 85638542, 264760, 264768, 264769, 264691. 35698423	264259, 66712502, 264682, 264683, 264635
kinase	UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	transcriptfactor	dna_ma_bind	synthase		- protease	UNCLASSIFIED
					Contains protein domain (PF00651) - transcriptfactor BTB/POZ domain	Contains protein domain (PF00096) - dna_ma_bind Zinc finger, C2H2 type		Contains protein domain (PF01852) - START domain	Contains protein domain (PF00090) - protease Thrombospondin type 1 domain	
Novel Protein sim. GBank gil5689407[db] BAA82987.1] - (AB028958) KIAA1035 protein [Homo saplens]	Novel Protein sim. GBank giļa836757]gbJAAD30541.1JAF13491 - (AF134918) semaphorin subclass 4 member G [Mus musculus]	95322893 (2635, 2636) Novel Protein sim. GBank gil4680204[gb]AAD27567.1[AF11417 - (AF114171) hypothetical protein [Sorghum bicolor]		Novel Protein sim. GBank giļ4240183 dbj BAĀ74870.1  - (AB020654) KIAA0847 protein [Homo sapiens]	86676351 (2641, 2642) Novel Protein sim. GBank gil4886505 emb CAB43377.1  - (AL050276) hypothetical protein [Homo sapiens]	87755272 (2643, 2644) Novel Protein sim. GBank gij5262591jemb CAB45736.1 - (AL080143) hypothetical protein (Homo saplens)	94845931 (2645, 2646) Novel Protein slm. GBank gil5459516 dbj BAA82407.1  - (AB029821) phosphatidylethanolamine N-methyltransferase [Homo sapiens]	87737614 (2647, 2648) Novel Protein sim. GBank gi 5031717 ref NP_005704.1 pGPBP - goodpasture antigen- START domain binding protein		87316289 (2651, 2652) Novel Protein slm. GBank gi[1397275 (U61947) - C06G3.8 gene product [Caenorhabdills elegans]
1316   85361609 (2631, 2632) Novel Protein sim. GBank (AB028958) KIAA1035 pro	88055167 (2633, 2634) Novel Protein sim. GBank gil4836757lgbJAAD30541. semaphorin subclass 4 mt	95322893 (2635, 2636) N 9 9 1	94238546 (2837, 2638)	86603567 (2639, 2640) Novel Protein sim. GBank (AB020654) KIAA0847 pre	86676351 (2641, 2642) N	87755272 (2643, 2644)	94845931 (2645, 2646)	87737614 (2847, 2648)	1325 94847471 (2649, 2650) Novel Protein sim. GBani the DPT/Kunitz family of factor pathway inhibitor p	87316289 (2651, 2652)
1316	1317	1318	1319	1320	1321	1322	1323	1324	1325	1326

Plant lipid transfer protein family 201531825, 29331822, 29331824, 29331827, 29331793, 29331793, 293373044, 8718518, 60432133, 22279000, 22279002, 284563, 264567, 264597, 264597, 264597, 264597, 264597, 264597, 264597, 264597, 264597, 264597, 264597, 264597, 264597, 264597, 264597, 264597, 26456	UNCLASSIFIED 264488, 22278997, 29331828, 264595, 18108351, 264766, 22279002, 264482, 264567			UNCLASSIFIED 264259, 29331826, 29331827, 35696052, 29331828, 60170831, 26448, 264686, 21906765, 55811957, 265020, 33657023, 33657109, 263973, 55811578, 35896423, 35695855, 56182323	UNCLASSIFIED 60432289, 66712502, 264591, 60433356, 60433438, 55812038, 265010, 264639, 55526488 lphosphalase 264905	ED
2_HUMAN - IIII ALU SUBFAMILY		Novel Protein sim. GBank gi 4678224 gb AAD26969.1JAC00713 - (AC007135) unknown protein [Arabidopsis thailana]	Novel Protein sim. GBank giļ437310 (L23504) - nodulin [Medicago truncatula]		Novel Protein sim. GBank gij5689527 dbj BAA83047.1  - (AB029018) KIAA1095 protein [Homo saplens] Novel Protein sim. GBank gil262535 (AF036685) - Similar	atase (Caenomabditis elegans) gi4240285[dbj]BA474921.1  - otein [Homo saplens]
100 100 100 100 100 100 100 100 100 100		87755276 (2657, 2658)			1333 88098476 (2665, 2666) Novel Protein sim. GBank (AB029018) KIAA1095 pro 1334 87592388 (2667, 2668) Novel Protein sim. GBank	to protein-tyrosine phosph 1335 87644798 (2669, 2670) Novel Protein sim. GBank (AB020705) KIAA0898 pr

264509, 264905, 264512, 264764, 264693, 264635, 264637	56182575, 56994075, 22278998, 22278999, 264092, 264259, 60432289, 29331826, 264306, 264309, 264112, 265008, 265009, 60433356, 55812038, 33657084, 265011, 265017, 265018, 265019, 264682, 26448, 264813, 264369, 265020, 264691, 27486261, 20281069, 18108379, 55811576, 35695655, 56182323, 60432113, 22279002, 264567	29331822, 265010, 264288, 264689, 18108370, 35695855	35696052, 264909, 264688, 264556, 264558	264505, 264507, 871065559, 264764 264681, 264685, 264686, 264692	264629	264910, 264686, 264534	263978	264909, 60170394	22278998, 264907, 264681, 264685, 264689, 265020, 264693, 22278000, 22279002, 264568	264906, 264908, 264909, 265006, 284910, 265011, 265017, 264764, 264765, 264767, 264769, 264631, 264634, 264638, 264567, 264486	264488, 264489, 264508, 264509, 264510, 264511, 264512, 264591, 264592, 264601, 264664, 264685, 264769, 264532, 264534, 264555, 264555, 264588, 22279002, 264486
	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED		tm7	struct		glycoprotein	kinase	glycoprotein	UNCLASSIFIED
					Contains protein domain (PF00001) - Im7 7 transmembrans receptor (rhodopsin family)	Contains protein domain (PF00560) - struct Leucine Rich Repeat			Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	Contains protein domain (PF00560) - glycoprotein Leucine Rich Repeat	
1336   87787890 (2671, 2672) Novel Protein sim. GBank gij465445[splP33465 VNUA_PRVKA - PROBABLE NUCLEAR ANTIGEN	94312042 (2673, 2674) Noval Protein sim. GBank gil5689471 dbj BAA83019.1  - (AB028990) KIAA1067 protein [Homo sapiens]		80249231 (2677, 2678) Novel Prolein sim. GBank gij1176422 (U43194) - mophilin [Mus musculus]		80089017 (2683, 2684) Novel Protein sim. GBank gil5019564 emb CAB44507.11 - (AL035542) d.1994E9.5 (hs6M1-17 (novel 7 transmembrane receptor (rhodpsin family) (olfactory receptor like) protein) [Homo saplens]	80082862 (2685, 2686) Novel Protein sim. GBank gil4557543 ref NP_001384.1 pECM2 - extracellular matrix protein 2		91225546 (2689, 2680) Novel Protein sim. GBank gi[2144101 pir  155210 -	80255717 (2691, 2692) Novel Protein sim. GBank gij3881052 emb CAA19523  - (AL023843) predicted using Genefinder; similar to seriner/hreonine kinase; cDNA EST yk248a12.3 comes from this gene; cDNA EST yk358c10.5 comes from this gene; cDNA EST EMBL:M89047 comes from this gene; cDNA EST yk246a12.5 comes	80417393 (2693, 2694) Novel Protein sim. GBank gil4504379[ref NP_003658.1 pHG38 - orphan G protein- coupled receptor HG38	87352335 (2695, 2696) Novet Protein sim. GBank gij3399720 dbj BAA32100  - (AB010999) pepiidylarginine deiminase type IV {Rattus norvegicus}
87787890 (2671, 2672) P	94312042 (2673, 2674)	80366114 (2675, 2676)	80249231 (2677, 2678)	88316311 (2679, 2680)	80089017 (2683, 2684)	80082862 (2685, 2686)	20562559 (2687, 2688)				
1336	1337	1338	1339	1340	1342	1343	1344	1345	1346	785	1348

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UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	dna_ma_bind
						Contains protein domain (PF00076) - dna_ma_bind RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)
1349   91225548 (2697, 2698) Novel Protein sim. GBank gi 2144101 pir  155210 - tricarboxylate carrier - rat (fragment)	(0	87361327 (2701, 2702) Novel Protein sim. GBank gil4887239 gb AAD32246.1  -   (AF064564) BAW protein [Fugu rubripes]	(4	1353   95345417 (2705, 2706) Novel Protein sim. GBank gil2144101 pir  155210 - tricarboxylate carrier - rat (fragment)	95350845 (2707, 2708) Novel Protein sim. GBank gil4689108lgblAAD27763.1lAF07703 • (AF077030) hypothetical 43.2 kDa protein [Homo saplens]	88260186 (2709, 2710) Novel Protein sim. GBank gij1469199 db  BAA09487  - (D50928) The KIAA0138 gene product is novel. [Ното sapiens]
91225548 (2697, 2698)		87361327 (2701, 2702)	80076386 (2703, 2704)	95345417 (2705, 2706)	95350845 (2707, 2706)	88260186 (2709, 2710
1349	1350	1351	1352	1353	35.	1355

					000000000000000000000000000000000000000
1356	1356   95313991 (2711, 2712) Novel Protein sim. GBank	Novel Protein sim. GBank gi 1113865 (U40342) - ninein Mare miscapies	<u>vs</u>	รเกต	18108387, <i>ZZZ1</i> 6995, <i>ZZZ1</i> 6996, <i>ZZZ1</i> 6996, 264094, 29331828, 264905, 265006, 265007,
				-	265008, 265010, 265017, 265018, 265019,
_				8	264764, 18108354, 264689, 21906765,
					265022, 18108364, 35696423, 83373044,
					18108387
1357	88260268 (2713, 2714)	88260268 (2713, 2714) Novel Protein sim. GBank gil897693 emb CAA90330  -	Contains protein domain (PF01852) -		264259, 29331822, 29331825, 264510,
		offine transfer protein [Bos taurus]	START domain	<u>~</u>	87168559, 265018, 264448, 264288,
			ſ	•••	21906765, 21906766, 21906768, 265021.
					264893, 18108376
1358	_	38719455 (2715, 2716) Novel Protein sim. GBank gil556219 (L36831) - transcription			264757
1359	1359 87771643 (2717, 2718)			UNCLASSIFIED	264907, 264909, 264510, 264511, 264512,
					18108331, 204704, 204334, 33031029,
					18108385, 264486, 264567
1360	87738272 (2719, 2720)	Novel Protein sim. GBank gi[2598282]emb CAA75612] -	S	synthase	60432289, 264605
		(Y15417) acetateCoA ligase [Coprinus cinereus]			
1361		05.1 -	Contains protein domain (PF00538) - UNCLASSIFIED		35696286, 22278997, 22278999, 264259,
			SAM domain (Sterile atpha motif)		29331826, 264508, 264509, 264905, 264907,
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					21906754, 87168474, 265011, 264761.
					264683, 264288, 264768, 264769, 264689,
					21906768, 265020, 265021, 33657023,
			`		55811576, 35696423, 264634, 60432113,
					22279002, 264482, 264486
1362	95287961 (2723, 2724)	95287961 (2723, 2724) Novel Protein sim. GBank gij5689411 dbij BAA82989.1  -	Contains protein domain (PF00400) - eph		56182575, 56181686, 60432049, 264259,
<u>.                                    </u>		(AB028960) KIAA1037 protein [Homo saplens]	WD domain, G-beta repeat		29331822, 56182181, 29331827, 35696052,
_					29331828, 264905, 264906, 264908, 264595,
					55812038, 85658542, 55811150, 264681,
					264288, 264369, 56181562, 60431528.
			,		55810764, 35696423, 60431850, 264558
1383	85758476 (2725, 2726)	1363   85758476 (2725, 2728) Novel Protein sim. GBank gil1130494 (U35776) - ADP-	Contains protein domain (PF01412) - UNCLASSIFIED	UNCLASSIFIED	264488, 29331826, 264907, 264687, 264689,
		ribosylation factor 1-directed GTPase activating protein	Putative GTP-ase activating protein		264693
		[Rattus norvegicus]	for Arf		
1364	88179488 (2727, 2728)	<del>-</del> -			60432289, 60433356, 60433438, 87168559,
	-				264603, 18108351, 21906766, 35698423,
					60432113
1365	-	83003108 (2729, 2730) Novel Protein sim. GBank gil4589562jdbjjBAA76803.11 -		oncogene	264766
	_	(AB023176) KIAA0959 protein (Homo sapiens)			
1366	87003262 (2731, 2732)	87003262 (2731, 2732) Novel Protein sim. GBank gil1084944 pir   S54495 -	Contains protein domain (PF00153) - transport	transport	265007
		hypothetical protein YPR021c - yeast (Saccharomyces	Mitochondrial carrier proteins		
		(Apiensiae)			

264488, 52646842, 52646365, 22278995, 56994075, 35696286, 22278998, 22278998, 22278998, 22278998, 22278998, 22278998, 22278998, 22278999, 264259, 264259, 35696052, 264905, 264907, 264509, 264909, 264900, 265000, 265000, 265000, 265000, 265000, 265001, 3657402, 55812038, 21908754, 265011, 3768559, 265017, 26488, 264787, 264689, 21908765, 21908766, 21908768, 21908768, 21908766, 21908768, 21908769, 55811957, 265020, 265021, 265022, 264534, 60170615, 264690, 264691, 18108352, 33657023, 33657349, 264628, 18108370, 18108374, 18108374, 56811576, 35696423, 35695855, 264635, 264556, 264537, 264556, 264537, 264558, 18108381, 18108385, 264555, 264537, 264556, 2645432, 60170394, 264558, 18108381, 264567	264259, 29331822, 29331826, 60432289, 29331827, 35696052, 264508, 284905, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264601, 264760, 264764, 264765, 264765, 264768, 2544629, 264699, 264634, 264634, 264637, 5264566, 264486, 264567, 833373044, 264563, 264566, 264486, 264567	265008, 60432229, 60433356, 33657084, 21906764, 21906769, 264555, 264638, 264559, 264638, 22278996, 264259, 29331822, 29331824, 29331826, 29331827, 35696052, 264508, 264509, 264907, 264908, 264907, 265011, 265019, 264764, 284766, 264768, 21906767, 35695917, 18108362, 35696423, 264632, 264632, 264638, 264632, 264638, 264552, 264638, 264552, 264638, 264552, 264638, 264552, 264638, 264552, 264638, 264552, 264638, 264552, 264638, 264552, 264638, 264552, 264638, 264552, 264638, 264552, 264638, 264552, 264638,	2227898, 264259, 29331822, 29331824, 22331825, 29331826, 29331827, 284905, 284509, 29331826, 29331827, 284905, 285009, 284788, 33557084, 85658542, 265010, 265018, 265019, 264762, 264448, 35695917, 33657109, 33657182, 33657182, 33657189, 33657182, 264563
UNCLASSIFIED .	оисовене	UNCLASSIFIED potassium_channel	glycoprotein
Contains protein domain (PF01342) - UNCLASSIFIED SAND domain	Contains protein domain (PF00071) -   oncogene Ras family		Contains protein domain (PF00335) - 4 transmembrane segments Integral membrane proteins
jij4884088jemb CAB43240.1  -	94320078 (2735, 2736) Novel Protein sim. GBank gi 464561 sp P35289 RB15_RAT RAS-RELATED PROTEIN RAB-15	86634033 (2737, 2738) Novel Protein sim. GBank gil2062702 (U90550) - butyrophilin [Homo sapiens] 95316910 (2739, 2740) Novel Protein sim. GBank gil5031823 refiNP_05823.1 pKCNM - potassium large conductance calcium-activated channel, subfamily M, beta member 2	95336512 (2741, 2742) Novel Protein sim. GBank gi 5032203 ref NP_005714.1 pTSPA - tetraspan 5
7 87721210 (2733, 2734)		1368   86634033 (2737, 2738) 1370   95316910 (2739, 2740)	1371 95336512 (2741, 2742)

263978	264769, 21906765, 21906767, 22278999, 264691, 264910, 55812038, 265010, 264681, 264684	264907, 264510, 265007, 264512, 265008, 60432229, 264689, 65274791, 264556, 264556, 23373044, 60432113	5264507, 52845158, 52646842, 52646365, 56182575, 56181686, 22278988, 56994075, 35696289, 22278988, 56994075, 35696289, 22278988, 56994075, 264259, 29331822, 52645080, 29331824, 29331825, 60432589, 29331824, 29331825, 60432289, 29331827, 264509, 264906, 264906, 264906, 264906, 264906, 264909, 265006, 264907, 264906, 29331830, 264909, 265006, 265017, 265018, 265019, 3109954, 3365702, 264448, 264764, 87168559, 264600, 265017, 265018, 265019, 55811150, 18108351, 264448, 264764, 264288, 264369, 264486, 2644229, 21906765, 21906766, 21906767, 21906768, 21906767, 21906769, 55811957, 35695917, 265020, 265031, 264630, 33657109, 27486261, 33657462, 264639, 264639, 264639, 264639, 264639, 264639, 264638, 18108336, 18108336, 18108336, 18108336, 264534, 264567, 22279000, 22279002, 264563, 264564, 264566, 284567	264768, 264769, 35695917, 22278997, 264691, 264259, 29331822, 264693, 35696052, 264693, 264508, 264509, 264906, 264628, 264508, 264629, 18108372, 264909, 264510, 284511, 264512, 265008, 264630, 264631, 264631, 264632, 264637, 264637, 264637, 264637, 264638, 26459, 264637, 264638, 264600, 264689, 265010, 265011, 264600, 264689, 264764, 264488, 264762, 264564, 264565, 264764, 264488, 264762, 264564, 264764, 264488, 264764, 264884, 264764, 264884, 264764, 264884, 264764, 264884, 264764, 264884, 264764, 264884, 264764, 264884, 264764, 264884, 264764, 264884, 26484, 264884, 26484,	gl 1663648 (U75321) - chromaffin Contains protein domain (PF00122) - ATPase_associated   29331824, 284591, 285019, 284686, 254706. [55811957, 284693, 22279002 [55811957, 284693, 22279002]
collagen	UNCLASSIFIED	Interferon	kinase	UNCLASSIFIED	- ATPase_associate
			Contains protein domain (PF00400) - kinase WD domain, G-beta repeat		Contains protein domain (PF00122)
80248517 (2743, 2744) Novel Protein sim. GBank gil840708 dbj BAA09334  -	r pagaraga agranda	95087036 (2747, 2748) Novel Protein sim. GBank gi[111876]pirj[JC1241 - beta- interferon-Induced protein - rat	94236942 (2749, 2750) Novel Protein sim. GBank gij5649176jgbjAAD03500.2  - (AF051155) G beta-like protein GBL [Rattus norvegicus] }		86864242 (2753, 2754) Novel Protein sim. GBank gij1663648 (U75321) - chromaffi granule ATPase II homolog [Mus musculus]
80248517 (2743, 2744)	80499421 (2745, 2746)				
1372	1373	1374	1375	1376	1377

22278994, 22278998, 264093, 264094, 264259, 29331824, 29331827, 265009, 26540, 4840835, 264288	265018, 265019, 16106351, 204359, 204259, 204269, 20446627, 25811957, 264691, 18108368, 264635, 263981, 18108385	265020	264510, 264512, 265009, 264288, 264564	87168559, 265017, 264628, 22279002	22278394, 22278397, 22278393, 23279223, 29331824, 29331826, 265007, 60432229,	60433356, 85658542, 265017, 265018, 264685, 264768, 21906766, 35695917, 33657023, 27486261, 27486262, 35695763, 35695855, 87168518, 22279002	18108396, 264692	264488, 264508, 264509, 264905, 264906.	264908, 224909, 264511, 264512, 284910, 264760, 18108351, 264766, 284769, 35695855, 264630, 264636, 264555, 264638, 264483, 264564, 264486	35696052, 55811386, 264688, 21906765, 265020, 33657023, 18108385	60432288, 29331828, 264908, 264907, 56182435, 265011, 264681, 60170815,	264591	2011520 02175000 051750 00331836	552/45/2, 222/0839, 204239, 2523/025. 29331827, 35696052, 264509, 264907,	33657402, 60433438, 264596, 21906754,	87168559, 264600, 265017, 264663,  18108354 52644229, 21906765, 21906766.	21906767, 21906768, 21906769, 265021,	264558, 60170394, 83373044, 22279000
			nuci_recpt	UNCLASSIFIED	UNCLASSIFIED		I INCI ASSIFIED	INCI ASSIFIED		UNCLASSIFIED	וען	UNCLASSIFIED		glycoprotein				
					Contains protein domain (PF00583) - UNCLASSIFIED									a.				
Novel Protein sim. GBank gil4107015 dbj BAA36293  - AB001772) PEM-5 (Ciona savignyi)		85679344 (2757, 2758) Novel Protein sim. GBank gij3252872 (AF035620) - BRCA1-	associated protein 2   profits   1980   87627962 (2759, 2760)   Novel Protein sim. GBank gil4837737 gb AAD30662.1  - (AF098834) germ cell specific Y-box binding protein [Homo	sapleins Novel Protein sim. GBank gil4718019919AD28508.11AF12538 - (AF125384) L82A Thoseoptia melanonasteri	,	•				86378788 (2769, 2770) Novel Protein sim. GBank gi[2384732 (AF015911) - NAC-1	91013049 (2771, 2772) Novel Protein sim. GBank gi[2384910 (AF022982) - contains similarity to the A-type potassium current class of	channel proteins [Ceenomabditis elegans]	(AJ132192) HS1 binding protein 3 [Mus musculus]	95101652 (2775, 2776) Novel Protein sim. GBank gil4895184[gb]AAD32753.1 AC00723 - (AC007231) putative	disease resistance protein [Arabidopsis thaliana]			
1378   87595071 (2755, 2756) Novel Protein sim. GBank gi (AB001772) PEM-5 (Clona s		85679344 (2757, 2758) N	87627962 (2759, 2760) N	88179656 (2761, 2762)	94847576 (2763, 2764)				86915895 (2767, 2768)	86378788 (2769, 2770)	91013049 (2771, 2772)	12 CT CT 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0		95101652 (2775, 2776)	. —			
1378		1379	1380	1381	1382			1383	1384	1385	1386		38	1388				

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Г						•			pentidase					_																											UNCLASSIFIED		LINCLASSIFIED	I INC. ASSIEIED	ONCEASOR LED
Contains protein domain (PF00841) - UNCLASSIFIED	Zn-finger in Ran binding protein and	others.								•									•	Contains protein domain (PF00709) -	Adenylosuccinate synthetase																				Contains protein domain (PF00560) - UNCLASSIFIED	Leucine Rich Repeat			
150000014LibA A B 2077 41	(poosso / judijować s / / ) -			•						94111916 (2779, 2780) Novel Protein sim. GBank gij3702295 (ACU05/83) -	R33083_1 [Homo sapiens]									Avel Protein sim GBank	SIN 3460101501028650101141 MOUSE .	ADENYLOSUCCINATE SYNTHETASE, MUSCLE	ISOZYME (IMPASPARTATE LIGASE)	lovel Protein sim. GBank gij726286 (U22394) - mSin3A	Mus musculus]																				95361471 (2789, 2790) Novel Protein sim. GBank gil2274845 dbj BAA21534  -  (DB8461) N-WASP [Rattus rattus]
	1389   91256016 (2777, 2778)   Novel Protein Sim. GBank gl						-	,		94111916 (2779, 2780) No	R							-		O422724E (2784 2782) Novel Protein sim GBank	(2012,1012) CC (221C)	ā <del>V</del>	31	94311097 (2783, 2784) Novel Protein sim. GBank					_												_	80409472 (2785, 2786)	-		95361471 (2789, 2790)
	1389 		_			_				1390										,	62			1392	_										_						[	1393		1394	1395

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	UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED  UNCLASSIFIED  Igf	
				Contains protein domain (PF00017) - UNCLASSIFIED Src homology domain 2  Contains protein domain (PF00790) - UNCLASSIFIED VHS domain Contains protein domain (PF00008) - tgf  EGF-like domain	
1396 95363253 (2791, 2792) Novel Protein sim. GBank gi[2135904[pit]  54810 - pHL E1F1 - human		91233667 (2795, 2796) Novel Protein sim. GBank gil5420389 emb CAB46680.1	87631076 (2797, 2798) Novel Protein sim. GBank gi 2496887[sp Q09232 YQ22_CAEEL - HYPOTHETICAL 32.0 KD PROTEIN C09F5.2 IN CHROMOSOME III	1400 95419084 (2799, 2800) Novel Protein sim. GBank gil283920 pir  S27939 - tensin - chlicken  1401 91226379 (2801, 2802) Novel Protein sim. GBank gil3256185 emb CAA15485  - (AL008635) dJ510H16.1 [Homo sapiens]  1402 95361475 (2803, 2804) Novel Protein sim. GBank gil1515427 (U57523) - nel homolog [Homo sapiens]  1403 94147933 (2805, 2806) Novel Protein sim. GBank gil5262615 emb CAB45747.1  - (AL080156) hypothetical protein [Homo sapiens]	
95363253 (2791, 2792)	87631317 (2793, 2794)	91233667 (2795, 2796)	87631076 (2797, 2798)	95419064 (2799, 2800) 91226379 (2801, 2802) 95361475 (2803, 2804) 94147933 (2805, 2806)	
1396			1399	1400 1400 1400 1400 1400 1400 1400 1400	

15	ווייים טיפני פסטסטים	1405 DESOCOSO 2000 2010 [Name Protein eim GBack nit 54065 jamh CAA583371.			264488, 56994075, 35696286, 29331822,
~	10103, 2010)	(X83413) U88 [Human beroesvitus 6]	, <del></del>		29331824, 29331826, 29331828, 35696052,
_					264508, 264908, 264907, 264908, 264510.
					264511, 264910, 33657402, 264594, 264758,
_					264600, 264604, 264762, 18108351, 264764.
					33657023, 33657109, 264628, 264634,
					83373044, 22279002, 264563, 264482.
					264486, 264567
8076	87847360 (7811 2812)	97612360 (2811 2812) Novel Protein sim GBank gil624076/gblAAC96425.11 -	8	collagen	264907, 264605
	01012303 (2011), 2012)	A 14260 Contains Dro deb Dy motife: SPKPP (20X)			
		(O42350) Contains Pro-ficial PA mones. Or No. (2007).			
		TETTA (3A), Sitting to solution portion cent wan process.			
		Corresponds to Swiss-Prot Accession Number P 13883			
┰	1100 0100 01000110	Neigh Control of the Court Alloyand Villas (AE015037)		UNCLASSIFIED	35696286, 22278999, 264094, 264259,
1407	94129872 (2813, 2814)	941298/2 (2813, 2814) Novel Protein Sim. Coank gilzoz 1000 (Artolous / 1			66714117, 29331826, 29331827, 29331828,
		endooligopeptidase A related protein; EUPA related protein			29146498, 264107, 264908, 265006, 265008,
_		[Orycolagus curiculus]			264910, 60433438, 265011, 265017,
_					18108351, 264448, 264288, 264686,
_					21906765, 21906769, 264692, 33657109,
_					18108370 264628 263972, 18108374,
_				•	35696423, 55811576, 264631, 264557,
_					200001101 20001101 1 1 1 1 1 1 1 1 1 1 1
_					264558, 83373044, 10106565, 67106319,
					60432113, 22279002
1408	95361477 (2815, 2816)	95361477 (2815, 2816) Novel Protein sim. GBank gi 2564953 (AF030001) -	Contains protein domain (PF00008) - oncogene	oncogene	264488, 264489, 35696286, 264109, 264508,
		unknown [Mus musculus]	EGF-like domain		2649US, 2645US, 2649US, 2649UT, 2649US,
					264909, 265008, 265009, 264910, 3365/402.
					264757, 264758, 265011, 264601, 265017,
					264760, 264762, 264683, 264685, 264766,
					264687, 264689, 21906767, 265021, 264690,
_				,	264691, 33657023, 264692, 264693,
					33657109, 264628, 264629, 35696423,
					35695855, 264631, 264632, 264634, 264635,
					264555, 264636, 264637, 264638, 56182323,
					264639, 264563, 264564, 264565, 264566.
					264567
1409	66644385 (2817, 2818)	86644385 (2817, 2818) Novel Protein sim. GBank git2662165(dbj]BAA23714  -			264693
		(AB007902) HH0712 cDNA clone for KIAA0442 has a 574-			
		be insertion at position 1474 of the sequence of KIAA0442.			
		[Homo sapiens]			
1410		86612587 (2819, 2820) Novel Protein sim. GBank	Contains protein domain (PF00386) -	complement	29331826, 264112, 264512, 265009, 265010,
_		gi[2493790[sp[Q60994]ACR3_MOUSE - 30 KD	C1q domain		264601, 264686, 264789, 21906767, 253974,
_		ADIPOCYTE COMPLEMENT-RELATED PROTEIN			284631, 284568
	-	PRECURSOR (ACRP30) (ADIPOCYTE SPECIFIC			
		PROTEIN ADIPODI			
١					

22278995, 22278997, 22278999, 29331822. 60432289, 29331828, 264907, 265017. 265019, 264682, 21906767, 21906768. 21906769, 265020, 264690, 264691. 33657023, 33657109, 27486264, 264628. 263972, 264634, 264558, 18108385	264757	56994075, 29331822, 35696052, 29331828, 29331830, 264809, 52644045, 264510, 52644296, 8568542, 81468414, 265017, 265018, 264681, 264687, 21908768, 35695917, 265020, 52644150, 264692, 264047, 2748624, 3569578, 2646870	18108387, 264566	264682, 264683, 265022, 264636	52646365, 56182575, 22278994, 22278995,	56994075, 22278996, 22278997, 22278998,	22278999, 264259, 29331822, 29331824, 29331825, 29331826, 29331827, 29331828	29146498, 66712502, 29331830, 52644045.	264113, 264511, 33657402, 264757,	21906754, 55811386, 265017, 265018,	265019, 264761, 264683, 264369, 264288,	264686, 264689, 21906766, 21906767.	29148627, 21906769, 55811957, 265020.	265021, 264690, 33657023, 65274620,	52645129, 27486262, 27486284, 60431528.	264629, 35695855, 56182323, 264559,	60432113, 264404, 22279002, 264482
kinasereceptor	UNCLASSIFIED	homeobox		UNCLASSIFIED		-											
Contains protein domain (PF00400) - (kinasereceptor WD domain, G-beta repeat		Contains protein domain (PF00023) - homeobox Ank repeat		Contains protein domain (PF00560) - UNCLASSIFIED Leucine Rich Repeat													,
1411   87818641 (2821, 2822) Novel Protein sim. GBank gij3123155 sp P91343 yW3M_CAEEL - HYPOTHETICAL 49.0 KD TRP-ASP REPEATS CONTAINING PROTEIN F55F8.5 IN CHROMOSOME I		1413 95416559 (2825, 2826) Novel Protein sim. GBank gij3879121 emb CAA94370j - (Z70310) predicted using Genefinder; Similarity to Mouse ankyrin (PIR Acc. No. S37771); cDNA EST EMBL:101923 comes from this gene; cDNA EST EMBL:D32335 comes from this gene; cDNA EST EMBL:D32335 comes from this gene; cDNA EST EMBL:D32723 comes from this gene; cDNA EST EMBL:D32723 comes from this		1414   94675860 (2827, 2828) Novel Protein sim. GBank gij3252981 (AF068921) - Ras- binding protein SUR-8 [Mus musculus]	94326948 (2829, 2830) Novel Protein sim. GBank gij1871187 (U90439) - unknown	protein [Arabidopsis thaliana]											
87818641 (2821, 2822)	84390919 (2823, 2824)	95416559 (2825, 2826)		94675860 (2827, 2828)													
1411	1412	1413		1414	1415												

18108392, 18108394, 18108397, 18108398, 22278995, 22278996, 35696286, 22278997, 22278998, 22278996, 25278997, 22278998, 22278999, 29331822, 35696052, 29331828, 29146498, 264905, 264907, 265007, 265007, 265008, 265007, 265007, 265011, 8718859, 265017, 265019, 264763, 264682, 26462, 18108351, 264681, 264682, 264682, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906766, 21906766, 21906768, 264689, 264634, 264634, 18108361, 33657023, 18108362, 18108379, 56811576, 65274791, 264634, 264634, 264634, 56182323, 18108381, 60170394, 18108385, 56526486, 87168518, 22279000	264107, 264448	264259, 264808, 255010, 52644229, 21906764, 21906768, 264690, 264639, 18108388	264259, 60432289, 265006, 87168474, 264288		18108358, 18108396, 18108397, 21906766, 18108398, 21906767, 56182575, 21908768, 21906768, 21906768, 21906769, 21906769, 21906769, 21906769, 21906769, 22278997, 22278997, 22278997, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 255021, 255022, 66170615, 29331824, 2484261, 29331828, 35695652, 29331828, 248907, 248907, 60431528, 66712502, 263972, 55811578, 35695423, 35695855, 264512, 265007, 60431850, 60432229, 60431735, 56182323, 264558, 18108385, 21906754, 55811386, 87168518, 87168558, 6043213, 265017, 265018, 265019, 22279002, 55811150, 264563, 264887, 18108391, 264763, 264486, 18108391
struct	ATPase_associated 264107, 264448	UNCLASSIFIED	struct	ATPase_associated	kinase
Contains protein domain (PF00735) - struct					
I - 1	87826663 (2833, 2834) Novel Protein sim. GBank gil4958935 dbj BAA78095.1  - (AB027570) suppressor of potassium transport defect 3 [Rattus norvegicus]	Ť	87757168 (2837, 2838) Novel Protein sim. GBank gi 2072294 (U95097) - mitotic phosphoprotein 43 [Xenopus laevis]	87298628 (2839, 2840) Novel Protein sim. GBank   gi 5174421 ref NP_008023.1 pCPNE - copine VI (neuronal)	94746986 (2841, 2842) Novel Protein sim. GBank gij3876090jembjCAA93459.1j - (Z69635) Similarity to Yeast uridine kinase (SW:URK1_YEAST); cDNA EST EMBL:214695 comes from this gene; cDNA EST CEMSE17F comes from this gene; cDNA EST yk209h1.5 comes from this gene; cDNA EST yk209h1.5 comes from this gene.
1416   84325977 (2831, 2832)   Novel Protein sim. GBank gil5106557 gb AAD39749, septin-like fusion protein [	87826663 (2833, 2834)	87594276 (2835, 2836)		87298628 (2839, 2840)	94746966 (2841, 2842)
1416	1417	1418	1419	1420	1421

				ſ	56004074 35606288 87168559 55811957
422 8	38178777 (2843, 2844)	1422   88178777 (2843, 2844) Novel Protein sim. GBank gil4505939 ref NP_000928.1 pPOLR - polymerase (RNA) II   DNA directed) polyneptide A (220kD)			55811578, 264555, 264557, 87168518
1423 8	86997762 (2845, 2846)			UNCLASSIFIED	264686, 264489, 264692, 264594, 264603, 265018, 264908
424	95201610 (2847, 2848)	1424 95201610 (2847, 2848) Novel Protein sim. GBank gij437181 (U02289) - GTPase- (activating prolein [Caenorhabditis elegans]	Contains protein domain (PF00620) - struct RhoGAP domain		29331622, 29331625, 29331627, 29146498, 264905, 264906, 264908, 264909, 265007, 264910, 265009, 33109954, 265010, 87168559, 265019, 264768, 264687, 21906766, 21906767, 21906767, 21906767, 21906767, 21906767, 21906767, 21906768, 2646891, 264692, 56526486, 22279002, 264691, 264692, 56526486, 22279002, 264683
1425	21662314 (2849, 2850)			UNCLASSIFIED	265007, 264558
1426	94322115 (2851, 2852)	94322115 (2851, 2852) Novel Protein sim. GBank gi 2078441 (U56964) - weak similarity to S. cerevisiae intracellular protein transport protein US)1 (SP:P25386) [Caenorhabditis elegans]		UNCLASSIFIED	2627898, 2527899, 264259, 60432049, 22278998, 22278999, 264259, 60432049, 26331827, 56182435, 264910, 60433356, 60433436, 21906756, 21906765, 21906765, 21906765, 21906765, 21906767, 21906769, 265020, 26502, 21906767, 18108370, 18108376, 26526486, 22279002, 264482
1427	91227510 (2853, 2854)	91227510 (2853, 2854) Novel Protein sim. GBank gij5816074(gblAADA5616.1)AF06194 - (AF061943) protate- derived STE20-like kinase PSK [Homo sapiens]	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	22278994, 56994075, 22278997, 29331828, 29331830, 264828, 265006, 265007, 255008, 265009, 264083, 264288, 18108354, 21906768, 29148629, 33657023, 18108374, 35695855, 83373044, 22278002, 284564
1428		94323008 (2855, 2856) Novel Protein sim. GBank gij138350jspjP28968jVGLX_HSVEB - GLYCOPROTEIN X PRECURSOR		glycoprotein	56181686, 264259, 264907, 265007, 265009, 264595, 265010, 264688, 65274620, 2646829, 6527479002, 264566
1429	87888689 (2857, 2858)			UNCLASSIFIED	264112, 264595, 265017, 265019, 21906763, 263977, 264555
1430	94735021 (2859, 2860)	94735021 (2859, 2860) Novel Protein slm. GBank gil1181619 db  BAA11565  - (D82364) a variant of TSC-22 [Gallus gallus]			264094, 29331824, 264591, 264593, 265018, 264681, 21906765, 21906767, 62774620, 55811576, 264639, 87168518, 22279002
1431		80429081 (2861, 2862) Novel Protein sim. GBank gij5420389jembjCAB46680.11 - (AJ243460) proteophosphogycan (Letshmania major)		UNCLASSIFIED	264768, 265020, 264691, 264693, 264638
1432		87463004 (2863, 2864) Novel Protein sim. GBank gil414797 (L18966) - pyruvate dehydronenase phosphatase [Bos taurus]		phosphatase	18108394, 29146496, 263007, 60433436, 264763, 29148629, 263969
1433		87605403 (2865, 2866) Novel Protein sim. GBank gil2460316 (AF022147) - uterus- overy specific putative transmembrane protein [Rattus	Contains protein domain (PF00100) - UNCLASSIFIED Zona pellucida-like domain	UNCLASSIFIED	264259, 264510, 264591, 264603, 264363
1434	85713730 (2867, 2868)	-		UNCLASSIFIED	264682, 264691

		(AB015330) HRIHFB2007 [Homo sapiens]			264259, 29331822, 29331824, 66714117, 29331826, 33656970, 264508, 264905, 66712502, 29331830, 264909, 265007, 265008, 264909, 265007, 265008, 264990, 265007, 265019, 265019, 18108351, 264762, 265017, 265018, 265029, 18108351, 264762, 21906768, 21906769, 21906767, 21906769, 21906769, 264691, 33657109, 264628, 18108374, 18108376, 5811576, 264638, 60170394, 56182323, 264559, 83373044, 87168518, 60452113, 22279000, 22279002,
1436	86635024 (2871, 2872) 87631082 (2873, 2874)	86635024 (2871, 2872) Novel Protein sim. GBank gij3183977 [emb[CAA39515] - (X56044) protein Htf9C [Mus musculus] 87631082 (2873, 2874) Novel Protein sim. GBank		UNCLASSIFIED	26-553, 264482, 264555 263978, 284557, 264559 22278997, 66714117, 29331826, 264907,
1438	85544280 (2875, 2876)		Contains protein domain (PF00595) - kinase PDZ domain (Also known as DHR or GLGF).	,	56182435, 265009, 18108351, 264692, 264693 264488, 264508, 264906, 264909, 264757, 264600, 264601, 264605, 264768, 264769, 264690, 35696423, 264558, 264563, 264568
1439		- A B	Contains protein domain (PF00450) - cathepsin Serine carboxypeptidase		264489, 18108394, 65274572, 56182575, 22278994, 22278996, 25278997, 22278998, 22278999, 264259, 60432049, 29331822, 29331824, 60432289, 29331827, 29331824, 60432289, 29331827, 29331824, 60432289, 264507, 264508, 264509, 264907, 66712502, 264509, 264907, 66712502, 265007, 265009, 60432229, 60433438, 265017, 265018, 265019, 18108354, 264289, 18108355, 264767, 21906765, 21906766, 21906769, 21906769, 25811957, 35895917, 265020, 265021, 265022, 33651109, 18108374, 265020, 264638, 56182323, 83373044, 18108387, 87168518, 60432113, 222790000, 264486
<u> </u>	87423643 (2879, 2880)	1440 87423643 (2879, 2880) Novel Protein sim. GBank gi 2662165 db  BAA23714  - (AB007902) HH0712 cDNA clone for KIAA0442 has a 574-bp insertion at position 1474 of the sequence of KIAA0442. [Homo sapiens]		UNCLASSIFIED	264887, 264259, 264906, 264907, 264908, 264909, 265008, 265010, 265017, 265018, 265019, 18108351, 264369, 265020, 33657023, 33657109, 60431528, 55811576, 264635

1441	95317662 (2881, 2882)	gij4493956jembjCAB11123.2j - lexExon; MAL3P6.28 protein, len: 167 aa; Similarity to ral proteins (C.elegans, lae & S.pombe). C.elegans	Contains protein domain (PF00646) - helicase F-box domain.	helicase	18108392, 264488, 263994, 264489. 5618255, 22278994, 22278995, 56994075, 35696286, 22278997, 22278999, 60432049, 264259, 29331822, 26331824, 56182181, 66714117, 29331822, 29331824, 56182181, 2645117, 29331826, 29331827, 29331828, 35696052, 29146498, 264508, 264908, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264609, 265017, 264604, 265018, 264609, 265019, 264761, 264761, 264761, 264609, 265019, 264761, 264609, 265019, 264761, 264609, 265019, 264761, 264609, 265019, 264609, 265019, 264609, 265019, 264609, 265019, 264609, 265019, 264609, 265019, 264609, 265019, 264609, 265019, 264609, 265019, 264609, 265019, 264609, 265019, 264609, 265019, 264609, 265019, 264609, 265019, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264637, 264639, 264639, 264637, 264639, 2646
1442	83367491 (2883, 2884)	Novel Protein sim. GBank gi 5103027 dbj BAA78765.1  - (AB023419) mSox7 (Mus musculus)		transcriptfactor	264567, 264488 264906, 265007, 264693, 264558
1443	87109935 (2885, 2886)		Contains protein domain (PF00435) - struct Spectrin repeat	struct	52645080, 264691, 264628, 264555
<del>4</del> 4	87620478 (2887, 2888)	Novel Protein sim. GBank gij3874447 emb CAB02772  - (281039) predicted using Genefinder; cDNA EST EMBL.:T01209 comes from this gene; cDNA EST yk278a11.3 comes from this gene; cDNA EST yk278a11.5 comes from this gene; cDNA EST yk308a9.3 comes from this gene; cDNA EST yk308a9.3 comes from this gene; cDNA EST yk308a9.5 com		UNCLASSIFIED	264259, 29331822, 29331824, 66714117, 29331828, 264288, 35695917, 33657023, 264635, 60170394
1445 1448		94990470 (2889, 2890) Novel Protein sim. GBank gil2959886jemb CAA11022  - (AJ222968) L-periaxin [Mus musculus] 85079068 (7801 2802)		UNCLASSIFIED	264369
1447	86945392 (2893, 2894)	Novel Protein sim. GBank gi 5081610 gb AAD39464.1 AF13544 - (AF135440) hunlington yeast partner C [Mus musculus]	Contains protein domain (PF01846) - FF domain		264369 18108398, 35696286, 22278997, 66714117, 29331828, 265009, 264758, 265018, 264288, 21906766, 21906767, 264692, 264634,
1448	94990477 (2895, 2898)	Novel Protein sim. GBank gij3980411 (AC004561) - putative proline-rich protein [Arabidopsis thaliana]	gi 3980411 (AC004561) - putative Contains protein domain (PF00439) - opsis thaliana] Bromodomain		29331827, 264509, 264909, 265008, 264595, 18108357, 18108385, 264566, 264486

66714117, 264906, 264908, 264591, 264601, 264764, 264632	35696286, 35696052, 265008, 265009, 60170831, 33109954, 264683, 264689, 35696423, 35695855, 56526486	29331826, 264683, 264693, 263978, 264630	264681, 33657023, 264629	264510, 264768	264556	29331822, 66714117, 29331825, 264905, 29331830, 265006, 265008, 265009, 265011,	265019, 18108351, 21906768, 33657109, 18108376, 264632, 56182323, 87168518	22278995, 22278996, 22278997, 22278998,	22278999, 29331827, 264107, 255017, 21906765, 21908766, 21908767, 21906769, 29148629, 18108370, 22278000	Γ	Π	29331826, 29331827, 29331828, 264102,	264107, 264110, 265009, 60432229, 265019, 265020, 265020, 265020, 265020, 264635, 26278002	264566	22278997, 264259, 29331824, 29331826,	29331827, 29331828, 265017, 265018,	264760, 264682, 264448, 264288, 264766,	265021, 264692, 3365/023, 3365/109, 35695855, 264566		22278997, 29331822, 35696052, 265009,	264758, 265017, 265018, 265019, 264760,	265022, 33657109, 27486261, 264555,	83373044	264555, 264556	60432049, 264259, 29146499, 264906, 264907 264419 265017 264766	18108270 18108274 284838 18108385
UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	peptidase	kinase		UNCLASSIFIED		transport		INCI ASSIFIED	transport	-			kinase				UNCLASSIFIED	gaba				UNCLASSIFIED	struct	
				,,,											Contains protein domain (PF00168) - kinase	C2 domain				Contains protein domain (PF00202) -	Aminotransferases class-III pyridoxal	phosphate			Contains protein domain (PF00385) - struct	Modified demain
	87458696 (2899, 2900) Novel Protein sim. GBank gij1707074 (U80450) - M01E11.2 [Caenorhabditis etegans]	87797970 (2901, 2902) Novel Protein sim. GBank gil4160304 emb CAA10600  - (AJ132192) HS1 binding protein 3 [Mus musculus]	85692899 (2903, 2904) Novel Protein sim. GBank gi[2832906]dbj[BAA24608.1] - (D89340) dipeptidvi peptidase III [Rattus norvegicus]	86130434 (2905, 2906) Novel Protein sim. GBank gij728831 sp P39188 ALU1_HUMAN - IIII ALU SUBFAMILY J WARNING ENTRY IIII	_			1456   86320218 (2911, 2912) Novel Protein sim. GBank	gij729230 spjP41004 CUT3_SCHPO - CHROMOSOME SEGREGATION PROTEIN CUT3		87800460 (2915, 2915) Novel Protein sim. GBank gil2246532 (U93872) - ORF 73	contains large complex repeal CR 73 [Kaposi's sarcoma-	associated herpesvirus]		95360920 (2917, 2918) Novel Protein sim. GBank	gij5524667igb AAD44333.1 AF15935 - (AF159356) Munc13-C2 domain	4 protein [Rattus norvegicus]			94741513 (2921, 2922) Novel Protein sim. GBank gil1707274 (U80931) - strong	similarity to class-III of pyridoxal-phoshate-dependent	aminofransferases [Caenorhabditis elegans]			88090605 (2925, 2926) Novel Protein sim. GBank gil1770466 emb CAA66912  -	(Kastas) M-pnase prosproprotein o Homo sapiens)
87860859 (2897, 2898)	87458696 (2899, 2900) I	87797970 (2901, 2902)   	85692899 (2903, 2904)	86130434 (2905, 2906)	_	1455 87797896 (2909, 2910)		86320218 (2911, 2912)		1800 C1001 0005000	87800460 (2915, 2916)								95354602 (2919, 2920)	94741513 (2921, 2922)						
1449	1450	1451	1452	1453	25	155		28		1467	Т				1459				1460	1461				462	1463	

264569, 22278995, 22278996, 22278997, 22278998, 29331622, 29331824, 29331825, 35696052, 20281100, 264905, 29331830, 264909, 265007, 33657402, 21906754, 265017, 285018, 264682, 264684, 264369, 264288, 264766, 21906765, 21906766, 21906767, 21906769, 35695817, 264691, 264630, 264631, 264659, 264651, 264651, 264651, 264651, 2646530, 2646539, 264565	264488, 22278994, 56994075, 60432049, 264259, 56182181, 60432289, 29331827, 52644045, 264511, 265007, 265008, 264596, 55812038, 55811386, 264600, 264602, 265017, 265018, 264604, 265019, 18108351, 18108354, 56181562, 21905769, 265021, 33657023, 33657182, 55811576, 264557, 18108382, 60432113	264512, 265017, 264689, 264558	264683, 264636	22278998, 264758, 265018, 265019, 21906769, 265020, 33657109, 22279002	18108394, 18108397, 18108398, 35696052, 29146499, 265007, 265008, 265009, 265010, 265011, 18108354, 18108365, 18108384, 18108388, 18108388	18108394, 18108398, 56182575, 264259, 29331822, 29331824, 29331825, 60432289, 264907, 264907, 264900, 265007, 264900, 265009, 264591, 60433229, 60433356, 264595, 60433438, 264758, 33109954, 265010, 265011, 265018, 264760, 264769, 265011, 265018, 264769, 265021, 18108351, 2906769, 55811957, 265021, 18108368, 264691, 18108362, 18108368, 264628, 18108378, 264637, 26457, 18108384, 18108388, 87168518, 60422113, 264404, 22279002, 264482, 264567, 264487	29331822, 29331824, 29331825, 264828, 264603, 264689, 264693, 18108374, 55811576
UNCLASSIFIED	glucoamylase	UNCLASSIFIED	struct		struct	UNCLASSIFIED	transport
							Contains protein domain (PF00153) - transport Mitochondrial carrier proteins
1464 87620482 (2927, 2928) Novel Protein sim. GBank gij3874447 emb CAB02772  - (Z81039) predicted using Genefinder; cDNA EST EMBL:T01209 comes from this gene; cDNA EST yk278a11.5 comes from this gene; cDNA EST yk278a11.5 tomes from this gene; cDNA EST yk308a9.3 comes from this gene; cDNA EST yk308a9.3 comes from this gene; cDNA EST yk308a9.5 com	87425192 (2929, 2930) Novel Protein sim. GBank gil4589596 dbj BAA76821.1  - (AB023194) KIAA0977 protein [Homo sapiens]		87614328 (2933, 2934) Novel Protein sim. GBank gi 2246532 (U93872) - ORF 73, contains large complex repeat CR 73 [Kaposi's sarcomalassociated herpesvinus]	95342862 (2935, 2936) Novel Protein slm. GBank gil4507241frefiNP_003137.1lpSSRP - structure specific recognition protein 1	79236174 (2937, 2938) Novel Protein sim. GBank gi 1906596 (U81788) - kinesin-73 [Drosophila melanogaster]	94990482 (2939, 2940) Novel Protein sim. GBank gil5649170lgblAAD43131.2lAF15909 - (AF159092) syld709613 protein [Homo sapiens]	87826842 (2941, 2942) Novel Protein sim. GBank gij3876146jembjCAB01750j - (Z78542) similar to Mitochondrial carrier proteins; cDNA EST EMBL:T01651 comes from this gene [Caenorhabditis elegans]
87620482 (2827, 2928)							
464	1465	1466	1467	1468	1469	1470	1471

94, 22278995, 56994075, 22278999, 22, 29331825, 22, 29331824, 66714117, 29331825, 26, 3565007, 264708, 265018, 265019, 264764, 264288, 264688, 24906769, 55811957, 285021, 264693, 264629, 55811578, 264634, 56182323, 22279002, 284566,	3, 35695917, 35696246, 264692, 723, 264693, 33657109, 35696052, 3, 264905, 264908, 264907, 264629, 3, 264909, 35696423, 35695855, 1, 264910, 264632, 264635, 5, 264637, 264556, 264567, 264599, 3, 60432113, 264604, 264685, 264766	1, 264682, 264288, 264568	179, 56161686, 25331624, 5045203, 826, 35696052, 264508, 264905, 264907, 264908, 264909, 264910, 33657402, 77, 264508, 265009, 264910, 33657402, 55, 264596, 55812038, 265011, 264691, 264685, 18, 264689, 55811957, 264691, 264692, 18, 18108370, 60431528, 18108374, 264635, 264636, 264536, 18108377, 264638, 264536, 264537, 264636, 264537, 26457, 264537, 264537, 264537, 264537, 264537, 264537, 264537, 264537	25, 264 107, 264 268, 265 0 20, 265 0 21, 74	98
					264686
Contains protein domain (PF00956) - N vicleosome assembly protein (NAP)	Contains protein domain (PF00400) - Is WD domain, G-beta repeat	Contains protein domain (PF00041) - Fibronectin type III domain	Conlains protein domain (PF00147) - Fibrinogen beta and gamma chains. C-terminal globular domain		
3688780 (AF042180) - testis- in [Mus musculus]	RO_BOVIN - CORONIN-LIKE		Novel Protein sim. GBank g 4757752 ret NP_004664.1 pANGP - angiopoietin 3	) Novel Protein sim. GBank gilz498308 sp Q60870 DP1_MOUSE - POLYPOSIS LOCUS PROTEIN 1 HOMOLOG (TB2 PROTEIN HOMOLOG) (GP108)	
87791609 (2845, 2846)	85800989 (2947, 2948)	75 86871935 (2949, 2950)		477 87774279 (2953, 2954)	1478 11754412 (2955 2956)
	1688780 (AF042180) - testis- Contains protein domain (PF00956) - MHC Nucleosome assembly protein (NAP)	protein Homo sapiens) 87791609 (2945, 2946) Novel Protein sim. GBank gij3688780 (AF042180) - testis- Contains protein domain (PF00956) - MHC specific Y-encoded-like protein [Mus musculus] Novel Protein sim. GBank gij3688780 (AF042180) - testis- Contains protein domain (PF00400) - struct gij2494890[sp]092176[CORO_BOVIN - CORONIN-LIKE WD domain, G-beta repeat PROTEIN P57	protein Homo sapiens)  87791609 (2945, 2946) Novel Protein sim. GBank gij3688780 (AF042180) - testis- specific Y-encoded-like protein [Mus musculus]  Nucleosome assembly protein (NAP)  Specific Y-encoded-like protein [Mus musculus]  Nucleosome assembly protein (NAP)  Specific Y-encoded-like protein [Mus musculus]  Nucleosome assembly protein (NAP)  Specific Y-encoded-like protein [Mus musculus]  Nucleosome assembly protein (NAP)  Specific Y-encoded-like protein [Mus musculus]  Nucleosome assembly protein (NAP)  Specific Y-encoded-like protein [Mus musculus]  Nucleosome assembly protein (NAP)  Struct  WD domain, G-beta repeat  PROTEIN P57  Contains protein domain (PF00041) - UNCLASSIFIED Fibroneckin type III domain	Protein Hone Sense 1988 Novel Protein sim. GBank See7191609 (2947, 2946) Novel Protein sim. GBank See7191609 (2947, 2946) Novel Protein sim. GBank See71935 (2949, 2950) See71, 2950) See71, 2950) See71, 2950 Novel Protein sim. GBank See71935 (2951, 2950) See71, 2950) See71, 2950 Novel Protein sim. GBank See71935 (2951, 2950) See71, 2950) See71, 2950 Novel Protein sim. GBank See71935 (2951, 2950) See71, 2950) See71, 2950 Novel Protein sim. GBank See71935 (2951, 2950) See71, 2950 Novel Protein sim. GBank See71935 (2951, 2950) See71935 (2951, 2950) See71935 Novel Protein sim. GBank See71935 (2951, 2950)	Specific Yencoded-like protein [Mus musculus]

					56182575 22278995 22278996 22278998
479	91640140 (2957, 2958)	1479   91640140 (2957, 2958) Novel Protein sim. GBank		pepudad	22278999, 29331822, 29331824, 66714117,
					264906, 264907, 56182435, 265006,
		kDa subunit [Homo sapiens]			60170831, 33657402, 264758, 33109954,
					21906754, 265017, 265019, 264448, 264288,
					264767, 264687, 52644229, Z1906764,
					264689, 21906/65, 21906/68, 21906/69,
-					265020, 265021, 60170615, 264691.
				•	33657023, 33657109, 33657162, 27466291,
					27486262, 33657349, 16106570, 60451526.
					263976, 55811576, 264556, 264557,
					60170394, 87168518, 264404, 22279000.
					22279UUZ, 204303, 204402
1480		94312412 (2959, 2960) Novel Protein sim. GBank gij3550456 emb CAA06329.1  -	5	UNCLASSIFIED	18108394, 652/45/2, 561625/5, 222/655.
		(AJ005073) Alix [Mus musculus]			230502500, 200524010, EEE, 2005; EEE, 2005
					222/0330, 222/0333, 204331, 20423;
					33383032, 28140483, 204103, 204109, 200100, 200100, 200100, 201100, 20
iii.					264108, 264907, 32644043, 204112, 203001.
					265008, 265009, 60433356, 60433438,
					264596, 33109954, 33657084, 5264429b,
	-				87168474, 285010, 87168559, 265017,
					265018, 265019, 264448, 264682, 264683,
					264769, 21906765, 21906766, 21908767.
					21908768 21906769 265020 265021.
					E1300105, E1300105, E300011, E1300105, E13001182.
	نى <u>د</u> .				251072 2460484 264557 263981
					2033/2, 50050005, E01001; E00001; 00033044 40400206 07460640 364666
					53373044, 16106363, 67166316, 203366,
	~		2	MHC	265006 265007 265010, 18108374
1481		87021442 (2961, 2962) Novel Prolein sim, GBank gil4836807]gbJAAD30566.1[AF14679 - (AF146793) PFT27		2	
3		Mus musculus	Contains protein domain (PF00008) - UNCLASSIFIED	JNCLASSIFIED	264908, 264910, 264758
1487		1JAF12292 - (AF122923) Wnt	EGF-like domain		
		Inhibitory factor-1 [Mus musculus]		1	284250 20131822 52645080 29331825
1483		Novel Protein sim. GBank gij535428 (U13736) - calmodulin-	Contains protein domain (PF00036) - S	iruci	29331826 33656970, 29331830, 265007,
_		like protein (Pisum sativum)	Er namu		55812038, 33109954, 265017, 264288,
					21906768, 21906769, 264636, 18108380.
					87168518, 22279000
1407		04131544 (2067 2058) Navel Protein eim GBank gil1911774 bhs1180090 -		UNCLASSIFIED	264489, 35696286, 264259, 264107, 264909.
<u> </u>					265008, 60433356, 33657402, 60433438,
		Ihuman, HeLa cells, Peptide Partial, 122 aal [Homo			264288, 21906765, 21906766, 29148627,
		saniens			33657023, 27486262, 18108374, 35696423,
					83373044, 60432113
1485	5 80194441 (2969, 2970	80194441 (2869, 2870) Novel Protein sim. GBank	Contains protein domain (PF00225) - struct	sluct	264369, 265020, 181083/4
_		gi[5360129]gb[AAD42883.1[AF15511 - (AF155117) NY-REN Kinesin motor domain	Kinesin motor domain		
		104 antigen [notitio bapteris]			

8	194125066 (2971, 2972)	1486 [94125066 (2971, 2972) Novel Protein sim. GBank gil4589516idbilBAA76780.11-	Contains protein domain (PF00069) - kinase		56182575, 22278999, 264906, 264907,
			Eukarvotic protein kinase domain		21906754, 87168474, 265017, 265019,
					18108351, 264288, 265020, 264566
1487	86452711 (2973, 2974)	86452711 (2973, 2974) Novel Protein sim. GBank gij5019275 embjCAB44431.1 - (AJ132751) xenablotic/medium-chain fatty acid:CoA ligase		synthase	21906754, 264486
		form XL-III [Bos taurus]	(677)		254686 254760 254680 284692 264693
<u>.</u>	87732026 (2975, 2976)	1488 87732026 (2975, 2976) Novel Protein sim. GBank	Contains protein domain (PF01443) - Igi		264509, 264708, 264907, 18108370, 264908,
		יייים (כניסדו ול) - פניסדו ולוי			264629, 264909, 264510, 265006, 264512,
					265007, 265008, 265009, 264555, 264556, 264557
1480		95104277 (2977 2978) Novel Protein sim GBank	Contains protein domain (PF00047) - prostaglandin		21906767, 22278999, 265022, 264259,
3		oii24973031spiQ62786iFPRP RAT - PROSTAGLANDIN F2- Immunoglobulin domain	Immunoglobulin domain		264693, 29331824, 29331825, 29331826,
		ALPHA RECEPTOR REGULATORY PROTEIN			29331827, 29331828, 264103, 263972,
		PRECURSOR (PROSTAGLANDIN F2-ALPHA RECEPTOR	:	•	66712502, 35696423, 35695855, 265007,
		ASSOCIATED PROTEIN)			265008, 265009, 83373044, 21906754,
					56526486, 265017, 264563, 18108351,
					264564, 264566, 264369, 264288
1490	87390127 (2979, 2980)			UNCLASSIFIED	56182575, 264259, 29331822, 29331824,
					66714117, 29331827, 29331828, 264508,
					264905, 66712502, 265007, 265008, 264594,
					33657402, 55812038, 87168474, 265018,
					18108351, 264369, 264288, 264769, 264689.
					21906767, 21906768, 55811957, 60170615,
					33657109, 35695855, 264635, 60170394,
					56526486, 22279002, 264563
1491		83594305 (2981, 2982) Novel Protein sim. GBank gi 295671 (L11275) - selected as		UNCLASSIFIED	265007, 264448, 18108372, 264558,
		a weak suppressor of a mutant of the subunit AC40 of DNA			56182323
		dependant RNA polymerase I and III (Saccharomyces			
ļ		Cerevisiae			
1492	85805363 (2983, 2984)	85805363 (2983, 2984) Novel Protein sim. GBank gi 1656005 (U71205) - rit [Mus musculus]	Contains protein domain (PF00071) - oncogene Ras family	oncogene	22278997, 22278998, 29331822, 264907, 66712502
ı					

		<del></del> T	T				Т		3	
26448B, 52646365, 65274572, 56182575, 22278994, 35696286, 56994075, 22278999, 60432049, 29331824, 29331828, 36696052, 26450B, 26450B, 264906, 526906, 5264045, 264909, 56182435, 265006, 265008, 265009, 60170831, 33657402, 55812038, 265010, 265011, 265011, 265018, 265019, 55811150, 264046, 2464682, 24906768, 21906768, 21906768, 21906769, 21906762, 21906761, 21906763, 21906764	264907, 265009, 264769, 18108370, 55811576, 264639, 264565, 264488	72278997, 264259, 264906, 264907, 265009, 264594, 33657084, 265017, 264760, 26448, 33657109, 264634, 56526486, 264563, 264568, 264565, 264566, 264488, 264567	35696286, 264906, 265019, 264693	264910, 264758, 265011, 264764, 264288, 264690, 264634, 264635, 56526486	264488, 65274572, 29331822, 265017, 265018, 21906765, 29148627, 29148629, 18108374, 264637, 264638, 264567	263978, 264566	22278999, 264769, 18108379	264559	264508, 264112, 264604, 264604, 32044130, 55811576, 264632, 284556, 264638, 56182323, 264563, 264486	29331822, 265007, 264369
SSIFIED		transcriptfactor	UNCLASSIFIED	kinase	oncogene	glucoamylase	UNCLASSIFIED	UNCLASSIFIED		- transferase
	Contains protein domain (PF01352) - kinase KRAB box	Contains protein domain (PF01352) - Iranscriptfactor KRAB box		Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain						Contains protein domain (PF00535) - transferase Glycosyl transferases
1493   91677215 (2985, 2986)   Novel Protein sim. GBank gij5689515 dbj BAA83041.1  - (AB029012) KIAA1089 protein [Homo sapiens]	2_HUMAN - IIII ALU SUBFAMILY	SB WAKNING EN KT !!!  87605267 (2989, 2990) Novel Protein sim. GBank gil4589588 dbj BAA76816.1  - (AB023189) KIAA0972 protein [Homo sapiens]	87784322 (2991, 2992) Novel Protein sim. GBank gil5420387 emb CAB46679.1 -	cDNA	90934938 (2995, 2996) Novel Protein sim. GBank gij728836 sp p39193JALU6_HUMAN - IIII ALU SUBFAMILY	SP WARNING ENTRY IIII 86451589 (2997, 2998) Novel Protein sim. GBank gi[2570198 (US4556) - microfilands sheath protein SHP3 [Litomosoides	sigmodonus	85795297 (3001, 3002) Novel Protein sim. GBank gil2078483 (U43200) - antifreeze glycopeptide AFGP potyprotein precursor (Boreogadus		87012701 (3005, 3006) Novel Protein sim. GBank gij3900855 (AC004874) - similar to N-acetylgalactosaminyltransferase; similar to Q07537 (PID:g1171989) [Homo sapiens]
(677215 (2985, 2986) N	37605265 (2987, 2988)	37605267 (2989, 2990)	87784322 (2991, 2992)	81695428 (2993, 2994)	90934938 (2995, 2996)	86451589 (2997, 2998)	10000 386 00000 AND		80206141 (3003, 3004)	87012701 (3005, 3006)
91	1494 8	1495 8	1496	1497	1498	1499	9	1501	1502	1503

1504	79640051 (3007, 3008)		Contains protein domain (PF00023) - UNCLASSIFIED Ank repeat		264693
1505		4753775 emb CAB41970.1  -  domo sapiens	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	29331826, 35686052, 264509, 264906, 264907, 264908, 264908, 264909, 264511, 264910, 55812038, 264759, 264763, 264764, 264689, 35695917, 265022, 33657109, 18108374, 264631, 264638, 264538, 264638, 264638, 264638, 264566
1506			Contains protein domain (PF00018) - glycoprotein SH3 domain	glycoprotein	65274572, 56182575, 56994075, 22278997, 22278998, 22278999, 264091, 264092, 260432049, 264259, 264091, 264092, 260432049, 264259, 5264050, 29331822, 26331827, 264106, 29331830, 264908, 26182435, 264110, 264681, 18108354, 264369, 264687, 264681, 18108354, 264369, 21906765, 29148627, 21906768, 21906765, 29148627, 264867, 264869, 21906765, 29148629, 5264150, 33657023, 18108376, 65274791, 56182323, 264558, 284559, 18108385, 87168518, 60432113, 22279000, 264565
1507		83738250 (3013, 3014) Novel Protein sim. GBank gij5689513[dbj BAA83040.1  -  (AB029011) KIAA1088 protein [Homo sapiens]		helicase	264639
1508					264593
509	87318451 (3017, 3018)	87318451 (3017, 3018) Novel Protein sim. GBank gil5031975 ref NP_005875.1 pPAK4 - protein kinase related to S. cerevislae STE20, effector for Cdc42Hs	Contains protein domain (PF00069) - kinase   PPAK4 - protein kinase related   Eukaryotic protein kinase domain ctor for Cdc42Hs	kinase	264259, 29331822, 29331824, 29331828, 264591, 33109954, 264563
1510	95362643 (3019, 3020)	Novel Protein sim. GBank gil113161[spiP28614 ACOR_ALCEU - ACETOIN CATABOLISM REGULATORY PROTEIN		UNCLASSIFIED	264259, 29331822, 265007, 18108374, 264556
1511		88318073 (3021, 3022) Novel Protein sim. GBank gi 728831 sp P39188 AU1_HUMAN - IIII ALU SUBFAMILY J WARNING ENTRY IIII		UNCLASSIFIED	264259, 29331822, 29331824, 29331827, 264509, 264907, 264510, 264511, 265007, 264512, 265008, 87168559, 264288, 265022, 33657023, 35695855, 264637, 264638,
1512		95345390 (3023, 3024) Novel Protein sim. GBank gil4559353lgblAAD23014.1IAC00658 - (AC006585) putative RIO1/ZK632.3/MJ0444 family extragenic suppressor protein [Arabidopsis thaliana]	Contains protein domain (PF01163) - RIO1/ZK632.3/MJ0444 family		52645156, 18108396, 56994075, 60432289, 265006, 60433356, 60433438, 21906754, 87168474, 87168559, 265018, 264762, 264763, 264687, 21906765, 21906769, 27486262, 35695763, 18108374, 35696423, 264555, 18108385, 18108387, 18108388, 87168518, 284482
1513		87436228 (3025, 3026) Novel Protein sim. GBank gi 1330394 (U58761) - C01F1.6 gene product [Caenorhabditis elegans]			35896052, 264905, 264907, 264908, 264908, 264908, 264909, 264910, 264591, 264766, 264689, 264636

52644507, 52645156, 52646365, 52646842, 65274572, 22278994, 35698286, 56994075, 264259, 52645080, 29331822, 29331825, 35698052, 29331825, 295000, 60433318, 60433438, 55812038, 21600764, 526440317, 52644296, 87168474, 87168559, 264448, 52644296, 21900765, 21900765, 21900767, 21900766, 356920, 52644150, 33657023, 52645129, 33657103, 33657103, 32695763, 18108376, 35696423, 35695855, 52644332, 18108385, 18108385, 87168518, 60432113	265020, 264639	265008, 56182323, 22279002	264091, 18108370, 264404	66714117, 264508, 264509, 264905, 264510, 264910, 264591, 264595, 264288, 264766, 264769, 18108374, 264638, 264638, 264486	264569, 264489, 60432049, 265009, 33657402, 264596, 21906754, 265019, 264369, 21906768, 21906769, 264691, 65274620, 33657182, 27486261, 18108374, 264557, 264639, 87168518, 22279002	18108392, 60432049, 264259, 29331824, 265007, 60433356, 265010, 21906768, 264636	18108394, 264259, 66714117, 265011, 264603, 265019, 18108364, 35696423, 264557, 264558, 18108388	29331824, 265018, 265020, 265021	65274572, 21906768, 264693	264112, 21906754, 263974	264683, 264687, 264689, 264690, 264692, 264693
UNCLASSIFIED	ubiquitin	UNCLASSIFIED		tm7		UNCLASSIFIED	- synthase	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED
Contains protein domain (PF01163) - UNCLASSIFIED RIO1/ZK632.3/MJ0444 (amily							Contains protein domain (PF00483) - synthase Nucleotidyl transferase				
live	79163536 (3029, 3030) Novel Protein sim. GBank gij3879501[emb CAA87795] - (Z47812) similar to ubiquitin carboxyl-terminal hydrolase; cDNA EST EMBL:D33366 comes from this gene; cDNA EST EMBL:D33965 comes from this gene; cDNA EST EMBL:D33822 comes from this gene; cDNA EST EMBL:D33827 comes from this gene; cDNA EST EMBL:D33827 comes from this gene; cDNA EST	88073539 (3031, 3032) Novel Protein slm. GBank gil498015 (L27479) - X123 [Homo sapiens]	87793325 (3033, 3034) Novel Protein sim. GBank gij3415134 (AF082024) - Phyb1 [Pimpinella brachycarpa]	ᅜᅙᇎ	94328689 (3037, 3038) Novel Protein sim. GBank gil5262681 emb CAB45771.1  - (AL080198) hypothetical protein (Homo sapiens)	87592855 (3039, 3040) Novel Protein sim. GBank gil2662161 dbj BAA23712  - (AB007900) HH0452 cDNA clone for KIAA0440 has a 438-bp insertion at position 1711 of the sequence of KIAA0440. [Homo saplens]	86970696 (3041, 3042) Novel Protein sim. GBank gi 5052351 gb AAD38516.1 AF13542 - (AF135421) GDP- mannose pyrophosphorylase B (Homo sapiens)	-	91005151 (3045, 3046) Novel Protein sim. GBank gij3776567 (AC005388) - Strong similarity to F2187.33 gij2809264 from A. thaliana BAC gbjAC002560. EST gbjN65119 comes from this gene. [Arabidopsis thaliana]		87799867 (3049, 3050) Novel Protein sim. GBank  gil4759040 ref NP_004283.1 pRIN1 - ras inhibitor
95345392 (3027, 3028) P	79163536 (3029, 3030)	88073539 (3031, 3032)	87793325 (3033, 3034)	87350697 (3035, 3036)		87592855 (3039, 3040)	86970696 (3041, 3042)	78960687 (3043, 3044)		80203723 (3047, 3048)	_
1514	1515	1516	1517	1518	1519	1520	1521	1522	1523	1524	1525

56182575, 264259, 60432049, 29331822, 60432289, 264908, 66712502, 60433438, 87168559, 265017, 264288, 21906768, 21906769, 263977, 55811576, 56182323, 18108381	22278995, 22278997, 284259, 66712502. 264596, 265017, 265019, 264682, 284448, 264683, 264764, 264685, 264686, 21906765, 21906766, 21906767, 21906768, 21906769, 265022, 264693, 83373044, 18108385	284488, 263994, 56182575, 22278895, 35696286, 22278997, 264259, 29331822, 60433289, 264308, 264509, 264906, 264907, 264908, 264909, 52644045, 264760, 264448, 265764, 265769, 264766, 18108357, 264768, 264766, 265021, 265021, 265021, 265021, 265021, 265021, 265022, 52644150, 33657109, 264629, 3569585, 60432113, 22279002, 264659, 3569585, 60432113, 22279002, 264563, 264564, 264486, 264567	264488, 264489, 35696286, 29331825, 35696526, 256906, 264907, 264905, 264906, 264907, 264909, 264511, 264512, 264907, 264509, 264591, 264511, 264764, 264683, 264684, 264768, 264768, 18108357, 264769, 35695917, 264628, 264631, 264634, 264565, 264636, 264631, 264633, 264569, 264486, 264486, 264489, 35695877, 264404, 264563, 264566, 264488	56182575, 35696286, 264097, 264259, 29331827, 29331827, 29331827, 29331827, 29331827, 29331827, 29331827, 25696052, 264509, 56182435, 25811386, 264581, 264581, 264386, 264288, 264766, 264687, 55811957, 35695917, 33657023, 35695763, 55810764, 35696423, 55811576, 263981, 60170394, 56182323, 83373044, 60432113, 264566
	UNCLASSIFIED	UNCLASSIFIED		rinase
				Contains protein domain (PF00595) - PDZ domain (Also known as DHR or GLGF).
Novel Protein sim. GBank gi 2792496 (AF041107) - tulip 2 [Rattus norvegicus]				95419351 (3061, 3062) Novel Protein sim. GBank gi(1905874 (U90878) - carboxyl terminal LIM domain protein [Homo sapiens]
88262512 (3053, 3054)	1528 94130918 (3055, 3056)	1529 94120793 (3057, 3058)	1530 95012765 (3059, 3060)	1531   95419351 (3061, 3062)
	ank gil2792496 (AF041107) - tulip 2	88262512 (3053, 3054) Noval Protein sim. GBank gil2792496 (AF041107) - tulip 2 [Rattus norvegicus] 94130918 (3055, 3056)	88262512 (3053, 3054) Novel Protein sim. GBank gil/406663 gb AAD20053  - tulip 2  [Rattus norvegicus] [Rattus norvegicus] [Rattus norvegicus]  94130918 (3055, 3056) [Rattus norvegicus]  64130918 (3055, 3056) Novel Protein sim. GBank gil/406663 gb AAD20053  - (AF131826) Unknown [Homo sapiens]	84130918 (3055, 3059) Novel Protein sim. GBank gilz792496 (AF041107) - fulip 2 [Rattus norvegicus] [AF130918 (3055, 3056)] [AF131826) Unknown [Homo sapiens]

22278994, 22278996, 35696286, 56994075, 22278997, 22278998, 22278999, 60432049, 22278999, 60432049, 22278999, 60432049, 22278997, 22278998, 22278999, 60432049, 28331825, 28331825, 6043289, 28331825, 28331827, 28331826, 23331826, 28331827, 285009, 33657402, 60433356, 60433438, 558412038, 24806754, 87168559, 265017, 265018, 265019, 18108351, 265020, 265021, 52644150, 33657023, 33657109, 27466261, 18108370, 35696423, 55811576, 65274791, 264558, 6618233, 60170394, 83373044, 87168518, 60432113, 22279000, 222779007, 264566	264369, 264691, 263978	264488, 264489, 22278999, 264259, 29331822, 35696052, 264508, 264509, 264905, 264908, 264511, 264512, 264905, 264901, 264907, 264908, 264761, 264762, 264448, 26476, 264588, 264687, 21906769, 55811957, 35695917, 265020, 264691, 33657023, 264692, 33657109, 264691, 18108374, 264632, 264584, 264565, 264566, 264486	18108392, 18108394, 65274572, 29331822, 264508, 265007, 265008, 265009, 265011, 264682, 18108354, 18108355, 52644150, 18108368, 264636, 18108381, 18108382	264106, 33109954, 265019, 264683, 35695917, 264690, 264692, 33657109	35686286, 56994075, 22278999, 35686052, 60433356, 60433438, 265011, 264683, 33657109, 35698423, 264631, 87168518, 22279000		П	264259, 29331822, 66714117, 265007, 55811386, 265010, 264600, 265017, 265019, 2642768, 264768, 265020, 265022, 55811576, 184108380, 264768, 265020, 265022, 55811576,
nuclease	UNCLASSIFIED	phosphatase	cathepsin	nuclease	nuci_recpt	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED
		·			Contains protein domain (PF00439) - nucl_recpt Bromodomain			
95337628 (3077, 3078) Novel Protein sim. GBank gij3218411jembjCAA19575.1j - (AL023859) SPBC19C7.07c, putalive tRNA splicing endonuclease ga mma subunit len:284aa, similar eg. to YAR008W, YAH8_YEAST, P39707, YAR008W, tRNA splicing endonuclease gamma subunit, (275aa), fasta scores, opt:289, E():6.4e-2	95352858 (3079, 3080) Novel Protein sim. GBank gi 5052634 gb AAD38647.1 AF14567 - (AF145672) BCDNA,GH12174 [Drosophila melanogaster]	Novel Protein sim. GBank gi 5052349 gb AAD38515.1{AF13501 - (AF135016) protein phosphatase 2A 48 kDa regulatory, subunit [Homo sapiens]	90937549 (3083, 3084) Novel Protein sim. GBank gi 5305702[gb AAD41779.1 AF12686 - (AF126867) calpain- like protease [Mus musculus]	84348768 (3085, 3086) Novel Protein sim. GBank gij728832 sp P39189 ALU2_HUMAN - !!!! ALU SUBFAMILY SB WARNING ENTRY !!!!	87757295 (3087, 3088) Novel Protein sim. GBank gij3493162 (AF084259) - bromodomain-containing protein BP75 [Mus musculus]	85757973 (3089, 3090) Novel Protein sim. GBank gi 1086591 (U41007) - similar to S. cervislae nuclear protein SNF2 (SP:P22082) in a a region of gly-arg repeats [Caenorhabdilis elegans]	Mount Dendoin nin Co.	[Mus musculus]
95337628 (3077, 3078)		95317848 (3081, 3082)		84348768 (3085, 3086)		85757973 (3089, 3090)	79475589 (3091, 3092) RE999594 (3093, 3094)	
1539	1540		1542	1543	1544	1545	1547	

94233065 (3095, 3096)	1548   94233065 (3095, 3098) Novel Protein sim. GBank gij3043692 dbj BAA25510  -		UNCLASSIFIED	29331824, 60431528, 264639, 56182323
330048 (3097, 3098)	95330048 (3097, 3098) Novel Protein sim. GBank gij5689519jdbjjBAA83043.1  - (AB029014) KIAA1091 protein [Homo saplens]		eph .	60424179, 22278995, 35696286, 22278998, 22278999, 264092, 264094, 29331822, 56182181, 29331824, 35686052, 264905, 264905, 264905, 264906, 264909, 265006, 264909, 265006, 264909, 265017, 265018, 265019, 55811386, 87168559, 265017, 265018, 264169, 5581150, 264682, 264288, 264369, 21906769, 25811957, 265020, 264981, 33657109, 60431528, 35696423, 35695855, 36526488, 60432113, 22279002, 264563, 264566
5201907 (3099, 3100)	95201907 (3099, 3100) Novel Protein sim. GBank gil544403lsplP35350jGUSB_BOVIN - POSSIBLE GUSTATORY RECEPTOR TYPE B (PPR1 PROTEIN)	Contains protein domain (PF00001) - Im7 7 transmembrane receptor (rhodopsin family)	m7	65274572, 60432289, 265008, 264910, 265011, 265017, 265019, 264768, 56182323
8077111 (3101, 3102)	pHS6S - heparan-sulfate 6-		UNCLASSIFIED	22278999, 29331822, 264508, 264509, 264906, 264907, 264907, 264909, 265007, 264512, 264910, 21906754, 265018, 265019, 264681, 264766, 264688, 264769, 21906769, 264692, 35695763, 264635, 264555, 264556, 264557, 264638, 264558, 264553
87617114 (3103, 3104)			UNCLASSIFIED	264259, 29331828, 66712502, 264764. 264288, 264688, 33657109, 264556
4725512 (3105, 3106)	94725512 (3105, 3106) Novel Protein sim. GBank gil4589570 dbj BAA76807.1  (AB023180) KIAA0963 protein [Homo sapiens]	Contains protein domain (PF00304) - dehydrogenase Gamma-thionins family	dehydrogenase	56182575, 35696286, 29146499, 264509, 264908, 264908, 264908, 56182435, 265006, 265008, 265008, 264909, 264757, 264758, 265017, 55811150, 18108351, 264764, 56181562, 35695917, 264693, 33657109, 18108374, 35696423, 65274791, 35695855, 264635, 264555, 56182323, 18108382, 83373044, 22279000
4233069 (3107, 3108)	94233069 (3107, 3108) Novel Protein slm. GBank gij3043692 dbjlBAA25510  - (AB011156) KiAA0584 protein [Homo saplens]	Contains protein domain (PF00446) - transferase Gonadotropin-releasing hormones	transferase	35696286, 22278997, 264259, 29331822, 29331824, 29331825, 265007, 265009, 60432229, 33657402, 55812038, 265011, 265019, 264681, 264369, 264767, 284768, 21906765, 21906769, 35695917, 264693, 18108370, 60431528, 55811576, 264631, 60170394, 56182323, 83373044, 18108385, 22279000, 22279002

264259, 29331826, 35696052, 264508, 264905, 264906, 264907, 264908, 52644045,	264909, 264910, 60432229, 50433356, 55812038, 264759, 33657084, 265812038, 264759, 33657084, 265011, 87168559, 264001, 285018, 265019, 264763, 264768, 264788, 264768, 264768, 264768, 35695817, 285022, 264691, 33657023, 35696423, 35695855, 264638, 264638, 264639, 18108385, 264555, 264638, 264638, 264639, 264638, 264639, 26463	83373044, 264758, 265022, 264600, 35696052, 264630, 35696423, 265011 264632, 264682, 29331822, 265020, 265011	264909, 264910, 264758, 55811386, 284761, 264909, 264910, 264762, 264690, 263978, 264762, 264690, 264986, 264692, 264639, 264690, 264998, 264639, 264699, 2646988, 2646988, 264698, 2646988, 2646988, 264698, 264698, 264698, 2646988, 264698, 264698, 264698,	22278999, 60432049, 264259, 29331824, 29331825, 29331825, 29331827, 264908, 264909, 60433365, 21906754, 265017, 265018, 264448, 21906767, 265021, 265022, 33657109, 18108370, 55811576, 83373044, 87188518, 22279000, 22279002	264259, 29331822, 60432289, 350956032, 264107, 264110, 21906754, 33109954, 87168559, 264760, 264763, 21906764, 21906765, 21906769, 265021, 264690. 35695855	264908, 264603, 264636	264510, 264594	264910 264764 264768	18108394 35698288 284259 29331822.	60432289, 35686052, 29331828, 264508, 66712502, 264908, 56182435, 265007, 264910, 60170831, 21906754, 265011, 265017, 265018, 265019, 18108351, 264448, 265021, 60170815, 264692, 35698423, 35695855, 264687, 56182323, 60432113, 22279002, 264482
INCLASSIFIED	•	UNCLASSIFIED	UNCLASSIFIED		glycoprotein	UNCLASSIFIED	03000	UNCLASSIFIED	debi-deposit	denya denya de
Contains protein domain (PF00400) - UNCLASSIFIED WD domain, G-beta repeat			Contains protein domain (PF00097) - UNCLASSIFIED Zinc finger, C3HC4 type (RING finger)		Contains protein domain (PF00560) - glycoprotein Leucine Rich Repeat				17700000	Contains protein domain (Pr U0441) - denydrogenase Acyl-CoA dehydrogenase
1555 87332970 (3109, 3110) Novel Protein sim. GBank gil2257495 dbj BAA21392  -   C			jij3329611 (AF078783) - 4-type zinc fingers (Pfam; zf- j; most similar to drosophila 33) (Caenorhabditis elegans)	94840376 (3115, 3116) Novel Protein sim. GBank gij5360105[gbJAAD42871.1]AF15510 - (AF155105) putative zinc finger protein NY-REN-34 antigen [Homo sapiens]	88224865 (3117, 3118) Novel Protein sim. GBank gij112908 sp P02730 A2GL_HUMAN - LEUCINE-RICH ALPHA-2-GLYCOPROTEIN (LRG)	84580675 (3119, 3120) Novel Protein sim. GBank gij3880146jemb[CAA92704] - (Z68319) Similarity to Human hnRNP F protein (PIR Acc. No. S43484); cDNA EST EMBL:D34218 comes from this gene; cDNA EST EMBL:D37248 comes from this gene; cDNA EST EMBL:D71817 comes from this gene; cDNA EST EMBL:D71817 comes from this gene; cDNA EST EMBL:D71817 comes from this gene; cDNA		(1)		87766371 (3127, 3128) Novel Protein sim. GBank gij1168287jspjP45953JACDV_RAT - ACYL-COA DEHYDROGENASE, VERY-LONG-CHAIN SPECIFIC PRECURSOR (VLCAD)
87332970 (3109, 3110)		1556 91228268 (3111, 3112)	1557 87640609 (3113, 3114)	1558 94840376 (3115, 3116)	1559 88224865 (3117, 3118)	1560 84580675 (3119, 3120)	1561 86609159 (3121, 3122)	Г	1583 85508694 (3125, 3126	1564 87766371 (3127, 3128

1565	87783381 (3129, 3130)	1565   87783381 (3129, 3130) Novel Protein sim. GBank gij129726;spjP05307jPDI_BOVIN - PROTEIN DISULFIDE ISOMERASE PRECURSOR (PDI) (PROLYL 4-HYDROXYLASE BETA SUBUNIT) (CELLULAR THYROID HORMONE BINDING PROTEIN) (P55)		ізотегазе	264488, 264689, 18108398, 55811957, 264534, 264259, 264508, 264905, 264509, 264908, 18108372, 264510, 264511, 264512, 265008, 264630, 265009, 264635, 264638, 264591, 264555, 264592, 264637, 264593, 264594, 264596, 265011, 264603, 22279002, 18108351, 264762, 264655, 264567
1566	87424749 (3131, 3132)	87424749 (3131, 3132) Novel Protein sim. GBank gij3880445 emb CAA20329  - (AL031268) VM105R.1 [Caenorhabdilis elegans]		tnf	22278996, 22278999, 264259, 29331822, 29331824, 60432289, 29331827, 66712502, 264908, 265008, 18108351, 52644229, 21906765, 21906767, 21906768, 21906769, 33657109, 264555, 264639, 264482
1567	84999006 (3133, 3134)	84999006 (3133, 3134) Novel Protein sim. GBank gil4929899jgbJAAD34110.1 AF15187 - (AF151873) CGI-115 protein [Homo sapiens]		UNCLASSIFIED	56182575, 21906769, 264692
1568	87648761 (3135, 3136)	87648761 (3135, 3136) Novel Protein sim. GBank gil4827063 refiNP_005072.1 pZNF1 - zinc finger protein 142 Zinc finger, C2H2 type (clone pHZ-49)	Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type		29331627, 29331830, 264511, 265009, 264758, 21906767, 21906768, 264691, 264693, 22279000, 22279002
1569	90936668 (3137, 3138)	90936668 (3137, 3138) Novel Protein sim. GBank gil5689451 dbjlBAA83009.1  - (AB028980) KIAA1057 protein [Homo saplens]	Contains protein domain (PF00443) - ubiquitin Ubiquitin carboxyl-terminal hydrolase family 2		65274572, 29331822, 29331824, 29331828, 264905, 56182435, 265007, 265019, 264764, 21906765, 21906769, 55811957, 60170615, 52644150, 264692, 33657023, 33657109, 18108377, 264563, 264567
1570		86943981 (3139, 3140) Novel Protein sim. GBank gi[1255430 (U53155) - No definition line found [Caenorhabditis elegans]		UNCLASSIFIED	264595, 264682, 265021
1571	T I	91210340 (3141, 3142) Novel Protein sim. GBank gil4507731[ref NP_001061.1[pTUBG - tubulin, gamma polypeptide	Contains protein domain (PF00091) - tubulin Tubulin/FtsZ famity	tubulin .	22278996, 35696286, 22278997, 264091, 264259, 29331824, 29331825, 29331827, 35698052, 264508, 264905, 56182435, 264510, 265007, 264788, 265011, 18108351, 264484, 264288, 264369, 21906765, 256021, 256021, 35657023, 264693, 18108377, 36996423, 35695855, 264555, 264555, 18108384
1572	81201664 (3143, 3144)			UNCLASSIFIED	35696052, 264905, 264906, 264908, 264910, 264758, 264766, 35695917, 264637
1573	1573 80207066 (3145, 3146)			UNCLASSIFIED	263972

							22278995, 29331822, 29331824, 29331826, 58182435, 284595, 55812038, 87168559, 265017, 264288, 21906764, 55811957, 35695917, 284692, 55811578, 264637, 5618233, 284559, 83373044, 60432113
ytochrome	phosphatase	UNCLASSIFIED	UNCLASSIFIED		UNCLASSIF	głycoprotein	
Contains protein domain (PF00173) - cytochrome Heme-binding domain in cytochrome b5 and oxidoreductases	Contains protein domain (PF00036) - phosphatase EF hand				Contains protein domain (PF00096) - UNCLASSIFIED Zinc finger, C2H2 type	Contains protein domain (PF00188) - glycoprotein SCP-like extracellular protein	
1574   94216142 (3147, 3148) Novel Protein sim. GBank gl 4758334 ref NP_004256.1 pFADS - delta-6 fatty acid desaturase	cDNA cDNA abditis	95314019 (3151, 3152) Novel Protein sim. GBank gi 2773195 (AF039711) - contains similarity to Physcomitrella patens glyceraldehyde 3-phosphate dehydrogenase (GB:X72381) [Caenorhabdilis elegans]	97613800 (3153, 3154) Novel Protein sim. GBank gi 2499130 sp P70315 WASP_MOUSE - WISKOTT- ALDRICH SYNDROME PROTEIN HOMOLOG (WASP)	-	88085141 (3157, 3158) Novel Protein sim. GBank gil2978255 dbjlBAA25190  - (AB007407) myeloid zinc finger protein-2 (Mus musculus)	87255702 (3159, 3160) Novel Protein slm. GBank gil4324682[gb AAD16986] - Contains protein domain (PFG (AF109674) late gestation lung protein 1 [Rattus norvegicus] SCP-like extracellular protein	95087431 (3161, 3162) Novel Protein sim. GBank gij2088838 (AF003386) - F59E12.4 gene product [Caenorhabditis elegans]
94216142 (3147, 3148)     (	95340019 (3149, 3150)	95314019 (3151, 3152)	87613800 (3153, 3154)	87123138 (3155, 3156)	88085141 (3157, 3158)	87255702 (3158, 3160)	95087431 (3161, 3162)
1574	1575	1576			1579		1581

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264259, 60432289, 29331827, 264509, 264905, 264906, 264907, 264909, 264910, 264762, 264288, 264768, 264769, 264632, 264555, 264639, 56528486, 22779000	357402, 264682, 21905/66. 1563	60424119, 52246842, 522/4517, 30102317, 22278995, 35696286, 22278999, 22278999, 264259, 29331822, 561218918, 22278999, 264259, 29331822, 56121891, 50424269, 60432289, 29331827, 29331828, 25640465, 56182435, 264510, 264451, 26182435, 264510, 264448, 264363, 265018, 18108358, 2564448, 264369, 265018, 265019, 264448, 264369, 265018, 265019, 264448, 264369, 265018, 265019, 264448, 264369, 265021, 265022, 29331822, 2559681, 264259, 29331822, 255021, 265022, 29331822, 255021, 265022, 29331822, 255021, 265022, 29331822, 255021, 265022, 29331822, 255022, 255022, 25502113, 22278002, 25502113, 22278022, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278022, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278022, 25502113, 22278002, 25502113, 22278002, 25502113, 22278002, 25502113, 22278022, 25502113,	29331824, 29331825, 264905, 265006, 265007, 265007, 265001, 265017, 26404, 26438356, 33109954, 87168474, 265011, 265017, 264004, 264389, 264685, 264769, 18108359, 263972, 18108383, 18108388, 264482, 263972, 18108383, 18108388, 264482, 264367, 264017, 264	264636, 264563 264636, 264563 264607, 264908, 264511, 264910, 264591, 264594, 284629, 264831, 264563, 264483,	264567 264259, 29331828, 264905, 265006, 264758, 21906754, 264761, 264762, 21906765, 21906769, 60170615, 52644150, 33657109, 35695855, 56182323, 18108385	65274572, 264490, 29331822, 66714117, 29331827, 56714117, 29331827, 29331828, 56182435, 265008, 60170831, 264595, 264758, 264586, 21906766, 21906768, 55811957, 27486265, 264639, 18108385, 56526486, 60432113
264259, 60432 264905, 26490 264762, 26428 264555, 26463	60170831, 336574( 35695855, 264563	60424179, 522 22278995, 356 22278999, 26, 60424269, 20, 35696052, 29 265008, 6043, 58 265008, 6043 264448, 27 27906765, 21 25695917, 26 35695917, 26 3569565, 56 3569565, 56	29331824, 26 265007, 2656 87168474, 20 264288, 264 21906765, 11 263972, 1811 264564	264636, 264563 264907, 264908 264907, 264908 264594, 264629	264259, 293 264259, 293 21906754, 2 21906769, 6 35695855, 6	65274572, 29331827, 29331827, 260170831, 264686, 21, 27486265, 60432113
				UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED
d.		Contains protein domain (PF00106) - dehydrogenase short chain dehydrogenase	Conains protein domain (rroudds) - priuspriatad		Contains protein domain (PF00468) - Ribosomal protein L34	
95358052 (3163, 3164) Novel Protein sim. GBank gil5420387 emb CAB46679.1  - (AJ243459) proteophosphoglycan [Leishmania major]	Novel Protein sim. GBank gi[5578958 emb CAB51351.1] -	531815lgbjAAD4482.11 -	87626117 (3169, 3170) Novel Protein sim. GBank gil4240132 dbj BAA74846.1  - (AB020630) KIAA0823 protein [Homo sapiens]	88057081 (3171, 3172) Novel Protein sim. GBank gij3785494 (AF098993) - No definition line found [Caenorhabditis elegans] 87617126 (3173, 3174) Novel Protein sim. GBank gij3253159 (AF005355) -	87802536 (3175, 3176) Novel Protein sim. GBank gil1077573 pir  S52680 - probable Contains protein domain (PF00468) - UNCLASSIFIED Ribosomal protein L34. mitochondrial - yeast (Saccharomyces cerevisiae)	90980653 (3177, 3178) Novel Protein sim. GBank gi[2137756 pir  148746 - semaphorin C - mouse (fragment)
5358052 (3163, 3164)	87622715 (3165, 3166)	<b>5337722 (3167, 3168)</b>	87626117 (3169, 3170) ,	88067081 (3171, 3172 87617126 (3173, 3174		
1582   93	583 8	1584	1585	1586 1587	1588	1589

5319	1590   95319825 (3179, 3180)		1	UNCLASSIFIED	264489, 22278996, 264259, 29331824,
					29331825, 29331826, 29331827, 265006,
			•		60433356, 21906754, 265017, 265018, 265019, 264448, 264784, 264288, 52644229
					21906765 21906767 21906768 21906769
					265021, 264692, 27486265, 35695763,
		-			56526486, 60432113, 22279000, 22279002
6877160	86877160 (3181, 3182)				264564
7882533	(3183, 3184)			MHC	264259, 264905, 29331830, 264595, 265017.
		gil4557749 ref NP_000237.1 pMHC2 - MHC class     transactivator			264448, 264288, 264690, 264629, 87168518
4991661	94991661 (3185, 3188)	_		UNCLASSIFIED	65274572, 60432049, 264509, 60433356,
					21906754, 21906767, 21906768, 18108370, 135606423, 22220000, 264665, 264667
7773752	(3187, 3188)	87773752 (3187, 3188) Novel Protein slm. GBank gij3877072jemb CAA87060  -		UNCLASSIFIED	264488, 29331827, 264905, 264906, 264907.
		(Z46937) similarity with ribosomal protein L21			264908, 264909, 264910, 264592, 264593,
		[Caenorhabditis elegans]			264757, 264602, 264604, 264760, 264681,
				•	264288, 264768, 264768, 29148629,
					35695917, 264692, 264628, 264629, 264630,
					264632, 264634, 264635, 264636, 264639, 264563, 264564, 264566
991942	5 (3189, 3190)	78919425 (3189, 3190) Novel Protein sim. GBank gil3152703 (AF065389) -	Contains protein domain (PF00335) - UNCLASSIFIED	UNCLASSIFIED	29331826, 264908, 55811957
		iens	4 transmembrane segments integral		
993392	79933928 (3191, 3192)			UNCLASSIFIED	29146498, 264758, 263967
697185	7 (3193, 3194)	86971857 (3193, 3194) Novel Protein sim. GBank	Contains protein domain (PF00067) - cyto450	cyto450	264092, 29331824, 264508, 264682, 264369.
		gi 5257114 gb AAD41244.1 AF09448 - (AF094480) cholesterol 24-hydroxylase [Homo sapiens]	Cytochrome P450		264686, 264630, 264563
786293	87862939 (3195, 3196)				264259, 264634
784982	9 (3197, 3198)	87849829 (3197, 3198) Novel Protein sim. GBank	Contains protein domain (PF01581) - UNCLASSIFIED	UNCLASSIFIED	52645080, 29331824, 29331826, 264511,
		gi[4506797 ref NP_000324.1 pSCA7 - spinocerebellar ataxia	.1pSCA7 - spinocerebellar ataxia FMRFamide related peptide family		265009, 265011, 264605, 264448, 264764,
					265020, 264692, 264693, 18108370, 264635, 18108385
1005600	80056002 (3199, 3200)				29331826, 264603, 264691, 264563
502324	15023248 (3201, 3202)			UNCLASSIFIED	264635
1692698	7 (3203, 3204)	86926987 (3203, 3204) Novel Protein sim. GBank	Contains protein domain (PF00018) - struct	struct	29146499, 264112, 264762, 18108351,
		gij5305704[gb]AAD41780.1 AF12853 - (AF128535) extoplasmic phosphoprotein PACSIN2 IMus musculusi	SH3 domain		29148627, 263974
1050207	2 (3205, 3206)	80502072 (3205, 3206) Novel Protein sim. GBank gi[283920 pir  S27939 - tensin -		collagen	264490, 29331824, 264907, 264909, 264511.
		chicken		,	265008, 264592, 265010, 265011, 264762,
					264764, 264369, 264288, 264687, 264769.
					264693, 264628, 264634, 264638, 264555,
					,264556, 264638, 264557, 264558, 264559, 18108385
3022181	3 (3207, 3208)			ATPase_associated	
		gi 4/68633 gb AAD29633.1 AF11682 - (AF116827)  unknown [Homo sapiens]			

CLASSIFIED CLASSIFIED CLASSIFIED					40.04	264905 264509 264906, 264907, 264908.
2012/103 (3211, 3212)   Novel Protein sim. GBank   Contains protein domain (PF00047) - struct   2   2   2   2   2   2   2   2   2	.~	91221129 (3209, 3210)				264909, 264604, 264766, 264768, 264692.
100   100						264693, 33657109, 264629, 35695855.
10871803 (3211, 3212)   Novel Protein sim. GBank   Gank Glavk   Contains protein domain (PF00047) - Struct   2						264635, 264636, 264637
10871805 (3213, 3214)   Novel Protein stim. GBank gil233920ipril[527939 - tensin - Gank gil234320ipril[527939 - tensin - Gank gil233920ipril[527939 - tensin - Gank gil23920ipril[527939 - tensin - Gank		94312703 (3211, 3212) N		Contains protein domain (PF00047) - s	truct	22278996, 22278999, 264259, 3365/402. 265017, 18108351, 26448, 21906767.
10871805 (3219, 3214) Novel Protein sim. GBank gil234520phi[527839 - tensin - chicken  94122843 (3217, 3218) Novel Protein sim. GBank gil239320phi[527839 - tensin - chicken  94122843 (3221, 3222) Novel Protein sim. GBank gil239320phi[527839 - tensin - chicken  94122843 (3221, 3222) Novel Protein sim. GBank gil239320phi[527839 - tensin - chicken  94122843 (3221, 3222) Novel Protein sim. GBank gil239320phi[527839 - tensin - chicken  94122843 (3221, 3222) Novel Protein sim. GBank gil23946pmb[CAA84337] - chicken  94122843 (3221, 3222) Novel Protein sim. GBank gil239446pmb[CAA84337] - chicken  94122843 (3221, 3222) Novel Protein sim. GBank gil23946pmb[CAA84337] - chicken  94122843 (3221, 3222) Novel Protein sim. GBank gil239446pmb[CAA84337] - chicken  94122843 (3221, 3222) Novel Protein sim. GBank gil239446pmb[CAA84337] - chicken  94122843 (3221, 3222) Novel Protein sim. GBank gil239446pmb[CA884337] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil239446pmb[CA884337] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil239446pmb[CA884337] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil239446pmb[CA884337] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil239446pmb[CA884337] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil239446pmb[CA884337] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil239446pmb[CA884304] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil239446pmb[CA884304] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil3874846pmb[CA884304] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil3874846pmb[CA884304] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil3874846pmb[CA884304] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil3874846pmb[CA884304] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil2394046pmb[CA884304] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil2394046pmb[CA884304] - chicken  185746031 (3223, 3224) Novel Protein sim. GBank gil2394	_	<u> </u>	1.1pMYOM - UNKNOWN			21906769, 52644150, 264691, 87168518
### ### ##############################		10871805 (3213, 3214) h	1 =		ranscriptiacion	600407
Monor Protein Sim. Gank gil224024113-11-   (AL049934) hypothetical protein [Homo sapiens]   (AL049934) hypothetical protein sim. GBank gil283920[pit][S27939 - tensin			promoted polypeptide		UNCLASSIFIED	264094, 264906, 264907, 264909, 264910,
			Novel Protein sim. Gbank gitzzałożajubjiprozobożi (AB002342) KIAA0344 [Homo sapiens]		-	264591, 264603, 26476B, 264693, 264034, 264635, 264637, 264639
94311572 (3217, 3218) Novel Protein sim. GBank gil283920[pril]527939 - tensin ;  (AL049934) hypothetical protein [Homo sapiens]  85468200 (3219, 3220) Novel Protein sim. GBank gil283920[pril]527939 - tensin ;  94122843 (3221, 3222) Novel Protein sim. GBank gil383920[pril]527939 - tensin ;  (Contains protein domain (PF00008) - peroxidase (PF111.17), thyrodia (Grave's disease) - human EGF-like domain (PF00008) - peroxidase (PF111.17), thyrodia (Grave's disease) - human EGF-like domain (PF00008) - peroxidase (PF00181) Signal (Bracycline residence) protein sim. GBank gil3874846[emb(CA494337] - (C10207) Smilarity to 8.subilis tracycline residence protein sim. GBank gil887486[emb(CA494337] - (C10207) Smilarity to 8.subilis tracycline residence protein sim. GBank gil887486[emb(CA494337] - (C10207) Smilarity to 8.subilis tracycline residence protein sim. GBank gil887486[emb(CA49337] - (C10207) Smilarity to 8.subilis tracycline residence protein sim. GBank gil887486[emb(CA49337] - (C10208) Smilarity to 8.subilis tracycline residence protein sim. GBank gil887486[emb(CA49337] - (C10208) Smilarity to 8.subilis (Grave's disease) - human EGF-like domain (PF00008) - (C10208) - (C10208						52644507, 52645156, 52646365, 52646842.
### (#################################			Novel Protein sim. GBank gil4884073 emb CAB43213.1  -	•		56182575, 22278994, 56994075, 35696286,
85468200 (3219, 3220) Novel Protein sim. GBank gil283920 ptr  527939 - tensin , chloken chloken			(ALCABSSA) Hypotherical profess (ALCABSSA)			22278997, 22278998, 22278999, 204293, 52645080, 29147620, 29331826, 35696052.
85468200 (3219, 3220) Novel Protein sim. GBank gij283920 pir  527939 - tensin ; chicken 94122843 (3221, 3222) Novel Protein sim. GBank gij107284 pir  A35415 - peroxidase (EC 1.11.17), thyroid (Grave's disease) - human EGF-like domain (fragment) 85746031 (3223, 3222) Novel Protein sim. GBank gij3874846 emb CAA94337  - (Z70307) Similarity to B. subtilis tetracycline resistance protein (SW:TCR2_BACSU); cDNA EST EMBL:C09951 comes from this gene; cDNA EST EMBL:C08265 comes from this gene (Caenorhabditis elegans)						33656970, 264508, 264509, 264907.
85468200 (3219, 3220) Novel Protein slm. GBank gil283920 pril[527839 - tensin ; chicken chicken gy122843 (3221, 3222) Novel Protein sim. GBank gil107284 pril[A35415 - contains protein domain (PF000008) - peroxidase peroxidase (EC 1.11.1.7), thyrold (Grave's disease) - human EGF-like domain (PF000008) - peroxidase (EC 1.11.1.7), thyrold (Grave's disease) - human EGF-like domain (F700008) - peroxidase peroxidase (EC 1.11.1.7), thyrold (Grave's disease) - human EGF-like domain (PF000008) - peroxidase peroxidase (EC 1.11.1.7), thyrold (Grave's disease) - human EGF-like domain (PF000008) - peroxidase peroxidase (EC 1.11.1.7), thyrold (Grave's disease) - human EGF-like domain (PF000008) - peroxidase peroxidase (EC 1.11.1.7), thyrold (Grave's disease) - human EGF-like domain (PF000008) - peroxidase peroxidase (EC 1.11.1.7), thyrold (Grave's disease) - human EGF-like domain (PF000008) - peroxidase (Froguential protein sin. GBank gil3874846 emb CAA94337  - (Z70307) Similarity to B.subdilis tetracycline resistance protein (SW:TCR2_BACSU); cDNA EST EMBL:C09951 comes from this gene: CDNA EST EMBL:C09265 comes from this gene: Caenom-baddilis elegans]						52644045, 56182435, 264510, 264511,
85468200 (3219, 3220) Novel Protein sim. GBank gij283920 pirij527939 - tensin - chicken chicken geroxidase (EC 1.11.7), thyroid (Grave's disease) - human EGF-like domain (PF00008) - peroxidase peroxidase (EC 1.11.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (F10007) Similarity to B.subtilis tetracycline resistance protein (SW:TCR2_BACSU); cDNA EST EMBL:C09951 comes from this gene; cDNA EST EMBL:C08265 comes from this gene (Caenonhabditis elegans) UNCLASSIFIED (UNCLASSIFIED)						264512, 3365/402, 21906/34, 32046511.  32106054 52644296, 87168474, 265017,
85468200 (3219, 3220) Novel Protein slm. GBank gij283920 prij 527939 - tensin -, chicken chicken gil283920 prij 527939 - tensin -, chicken geroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (PF00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.1.7), thyroid (Grave's disease) - human EGF-like domain (F00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (F000008) - peroxidase (EC 1.11.1.7), thyroid (EC 1.11.1.7), thyroid (EC 1.11.1.7),						265018, 265019, 18108351, 264448, 264288.
85468200 (3219, 3220) Novel Protein sim. GBank gi 283920 pir  S27939 - tensin -, chicken chicken chicken chicken chicken chicken chicken g4122843 (3221, 3222) Novel Protein sim. GBank gi 107284 pir  A35415 - Contains protein domain (PF00008) - peroxidase peroxidase (EC 1.11.7), thyrold (Grave's disease) - human EGF-like domain (fragment) (fragment) (fragment) (fragment) (fragment) (fragment) (270307) Similarily to B.subtilis tetracycline resistance protein (SW:TCR2_BACSU); cDNA EST EMBL:C09951 comes from this gene; cDNA EST EMBL:C09265 comes from this gene (Caenorhabditis elegans) UNCLASSIFIED (UNCLASSIFIED)				1		264769, 52644229, 21906765, 21906766,
85468200 (3219, 3220) Novel Protein slm. GBank gi 283920 ptr  S27939 - tensin ;  chicken  chicken  94122843 (3221, 3222) Novel Protein sim. GBank gi 107284 ptr  A35415 - Contains protein domain (PF00008) - peroxidase  peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain  (fragment)  (fragment)  (fragment)  (Z70307) Similarity to B.subtilis tetracycline resistance protein (SW:TCR2_BACSU); cDNA EST EMBL:C09951  comes from this gene; cDNA EST EMBL:C09265 comes  from this gene [Caenorhabditis elegans]  UNCLASSIFIED				j		21906767, 21906768, 21906769, 55011557, 21606767, 21906768, 21906769, 55022
85468200 (3219, 3220) Novet Protein slm. GBank gij283920 pir  S27939 - tensin ; chicken 94122843 (3221, 3222) Novel Protein sim. GBank gij107284 pir  A35415 - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (fragment) (fragment) (fragment) (270307) Similarity to Bank gij3874846 emb CAA94337  - (Z70307) Similarity to Baubtilis tetracycline resistance protein (SW:TCR2_BACSU); cDNA EST EMBL:C09951 comes from this gene; cDNA EST EMBL:C098265 comes from this gene (Caenorrhabditis elegans) UNCLASSIFIED						52644150, 33657023, 33657109, 52645129.
85468200 (3219, 3220) Novet Protein sim. GBank gij283920 pir  S27939 - tensin ; chlcken 94122843 (3221, 3222) Novet Protein sim. GBank gij107284 pir  A35415 - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (fragment) (fragment) (fragment) (fragment) (fragment) (fragment) (grave's disease) - human EGF-like domain (fragment) (fragment) (fragment) (fragment) (fragment) (grave's disease) - human EGF-like domain (fragment) (fragment) (fragment) (fragment) (fragment) (grave's disease) - human EGF-like domain (fragment) (f						27486261, 27486262, 35695763, 264628,
85468200 (3219, 3220) Novel Protein sim. GBank gi 283920 pir  S27939 - tensin - chicken chicken 94122843 (3221, 3222) Novel Protein sim. GBank gi 107284 pir  A35415 - Contains protein domain (PF00008) - peroxidase peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (fragment) (fragment) 85746031 (3223, 3224) Novel Protein sim. GBank gi 3874846 emb CAA94337  - (Z70307) Similarity to Baubtilis tetracycline resistance protein (SW:TCR2_BACSU); cDNA EST EMBL:C09951 comes from this gene; cDNA EST EMBL:C098265 comes from this gene [Caenorrhabditis elegans] UNCLASSIFIED						18108370, 18108376, 35696423, 264638,
85468200 (3219, 3220) Novel Protein sim. GBank gi[283920 pir][S27939 - tensin ; chicken chicken gli283920 pir][S27939 - tensin ; chicken gli283920 pir][S27939 - tensin ; chicken gli282843 (3221, 3222) Novel Protein sim. GBank gi[307284 pir][A35415 - Contains protein domain (PF00008) - peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (FF00008) - peroxidase (Fragment) (fragment) (Fragment) (A70307) Similarity to B.subiliis tetracycline resistance protein (SW:TCR2_BACSU); cDNA EST EMBL:C09951 comes from this gene; cDNA EST EMBL:C08265 comes from this gene [Caenorhabditis elegans] UNCLASSIFIED (UNCLASSIFIED)						52644332, 18108387, 87156318, 2278000. 264563, 264486
85468200 (3219, 3220) Nover Protein sim. Cbank git203920[pnt]527 503 - tensing chicken  94122843 (3221, 3222) Novel Protein sim. GBank git107284[pirt]A35415 - Contains protein domain (PF00008) - peroxidase peroxidase (EC 1.11.1.7), thyrold (Grave's disease) - human EGF-like domain (PF00008) - peroxidase (EC 1.11.1.7), thyrold (Grave's disease) - human EGF-like domain (FF00008) - peroxidase (EC 1.11.1.7), thyrold (Grave's disease) - human EGF-like domain (FF00008) - peroxidase (EC 1.11.1.7), thyrold (Grave's disease) - human EGF-like domain (FF00008) - peroxidase (Fromain in this density of Sank gil3874846]emb[CAA94337] - (Z70307) Similarity to B.subtilis tetracycline resistance protein (SW:TCR2_BACSU); cDNA EST EMBL:C09951	1	_	cipaci (COZCE)		UNCLASSIFIED	264593, 264757, 55812038, 265018, 265020,
94122843 (3221, 3222) Novel Protein sim. GBank gi 107284 pir  A35415 - Contains protein domain (PF00008) - peroxidase peroxidase (EC 1.11.1.7), thyroid (Grave's disease) - human EGF-like domain (fragment)  (fragment)  85746031 (3223, 3224) Novel Protein sim. GBank gi 3874846 emb CAA94337  - (Z70307) Similarity to Baubtilis tetracycline resistance protein (SW:TCR2_BACSU); cDNA EST EMBL:C09951 comes from this gene; cDNA EST EMBL:C09956 comes from this gene (Caenorhabditis elegans)	0		Novel Protein Sim. Gbank gijzosszuprijjsz rasa - tensin g chicken			264691, 264692, 264693, 264631, 264634, 264635, 264555, 22279000, 264564
94122843 (3221, 3224) Novel Protein Sim. Obains 9110120170  peroxidase (EC 1.11.17), thyroid (Grave's disease) - human EGF-like domain  peroxidase (EC 1.11.17), thyroid (Grave's disease) - human EGF-like domain  (fragment)  85746031 (3223, 3224) Novel Protein sim. GBank gij3874846jemb CAA94337  -  (270307) Similarity to B.subtilis tetracycline resistance protein (SW.TCR2_BACSU); cDNA EST EMBL:C09951 comes from this gene; cDNA EST EMBL:C08265 comes from this gene (Caenomabditis elegans)	- 1			Contains protein domain (PF00008) -	peroxidase	35696286, 21906765, 264691, 35696423
(fragment)   (fragment)   85746031 (3223, 3224)   Novel Protein sim. GBank gil3874846 emb CAA94337  -				in EGF-like domain		
85746031 (3223, 3224) Novel Protein sim. GBank gij38 4846jemb CAA94337/- (Z70307) Similarity to B.subilis tetracyclina resistance protein (SW:TCR2_BACSU); cDNA EST EMBL:C09951 comes from this gene; cDNA EST EMBL:C08265 comes from this gene [Caenorhabditis elegans]	- 1		(fragment)		UNCI ASSIFIED	264488, 264509, 18108370, 18108387.
(Z70307) Similarity to B.subtilis tetracycline resistance protein (SW:TCR2_BACSU); cDNA EST EMBL:C09951 comes from this gene; cDNA EST EMBL:C08265 comes from this gene [Caenorhabditis elegans]	10		Novel Protein sim. GBank gij3874846 emb CAA94337 -		-	264486
from this gene [Caenorhabditis elegans]			(Z70307) Similarity to B.subtilis tetracycline resistance			
from this gene [Caenorhabditis elegans]			Comes from this gene: cONA EST EMBL: C08265 comes			
UNCLASSIFIED			from this gene (Caenorhabditis elegans)			754750
	12	1 82247354 (1225 122R)	_		UNCLASSIFIED	264739

<u> </u>	1614 [91228634 (3227, 3228) Novel Protein sim. GBank			UNCLASSIFIED	22278995, 22278996, 22278997, 22278998,
_		[gi 4680673 gb AAD27726.1 AF13295 - (AF132951) CGI-17	eRF1-like proteins		22278999, 264259, 29331822, 264908,
		protein [Homo sapiens]			264512, 265009, 265011, 265017, 265018, 265019 18108351 284683 264288 264766
					21906767, 21906768, 21906769, 35695917,
					265021, 265022, 35696423, 35695855,
					60170394, 56182323, 83373044, 264566
	86121909 (3228, 3230)	86121909 (3228, 3230) Novel Protein sim. GBank gij5689485 dbj BAA83026.1  -	Contains protein domain (PF00023) - homeobox	нотворох	22278996, 35696286, 22278997, 29331822,
		(AB028997) KIAA1074 protein [Homo sapiens]	Ank repeat		35696052, 29331828, 264508, 264908,
		•			264909, 56182435, 264511, 265017, 265019.
					264766, 264767, 264768, 265020, 264691,
	·				264628, 264632, 264635, 264555, 264556,
					56182323, 264558, 22279002
	94311819 (3231, 3232,			UNCLASSIFIED	264488, 52644507, 52645156, 52646365,
		(Z78418) cDNA EST EMBL: D71020 comes from this gene;			52646842, 22278994, 22278995, 35696286,
		cDNA EST EMBL: D73593 comes from this gene; cDNA			22278996, 22278997, 22278999, 52645080,
	,	EST EMBL: C07649 comes from this gene; cDNA EST			29331822, 29331824, 29331825, 29331827,
-		EMBL:C09081 comes from this gene; cDNA EST yk399f2.3			29331828, 35696052, 33656970, 264905,
		comes from this gene; cDNA			264909, 264594, 52646317, 21906754,
_					33657084, 52644296, 87168474, 87168559,
					265017, 265018, 265019, 264681, 264448,
_					264684, 52644229, 21906784, 264689,
					21908765, 21906766, 21906769, 35695917,
					265020, 265021, 52644150, 33657023,
					52645129, 33657109, 33657182, 27486261,
					27486262, 33657349, 27486265, 35695763,
					18108376, 35696423, 35695855, 264557,
	-				52644332, 264558, 18108385, 87168518
	88090742 (3233, 3234		Contains protein domain (PF01529) - peptidase	peptidase	35696052, 264905, 264509, 264907, 264908,
		gil466053[splP34679[YO41_CAEEL - HYPOTHETICAL 68.7] DHHC zinc finger domain	7 DHHC zinc finger domain		264510, 264511, 264764, 264766, 264768,
		KD PROTEIN ZK757.1 IN CHROMOSOME III			264689, 264693, 18108374, 264635, 264636.
					264638
1618	_	86272860 (3235, 3236) Novel Protein sim. GBank gil4240231 dbjjBAA74894.11 -		struct	35696286, 22278999, 264092, 29331824,
		(ABUZU678) KIAAU871 protein [Homo sapiens]			29331825, 35696052, 33657084, 21906765,
	1619 95354580 (3237 3238	95354580 (3237 3238) Novel Protein sim CBank	Contains assists domain (DE00010)	transcription to the contraction of the contraction	6740647 66774677 22278000 264260
		dis031763ted0NP 005515 tloHRVI - hairy (Orosophia)	Holix-loop-holix DNA-hloding domain	ionalidi Della i	2010182 0211314, 2111033, 20123,
		homotoa			29331827 29331828 35896052 56182435
				•	265007, 265008, 264910, 60170831.
					E0432220 E043334 E0433434 265019
					264448 264288 264686 21906768 265021
					60170615 33657023 65274620 33657109.
					18108374 18108378, 35696423, 35695855,
					56182323, 56526486
	1620  87344655 (3239, 3240	87344655 (3239, 3240) Novel Protein sim. GBank  gi 1351047 sp P45843 SCRT_DROME - SCARLET		UNCLASSIFIED	264684
		PROTEIN			

LINCLASSIFIED 1264930	in domain (PF00622) - UNCLASSIFIED	ribosomalprot 18108398, 264259, 264809, 56182435, 87168474, 264448, 21906768, 35695917, 264691, 87168518, 264563	UNCLASSIFIED 68714117 29331825, 264909, 265008, 264758	kinase 264489, 22278994, 22278996, 22278996, 35696286, 22278994, 22278996, 22278999, 264092, 264259, 29331824, 29331825, 22331827, 29331827, 29331827, 29331828, 264108, 224508, 33657084, 265017, 265018, 18108351, 264683, 264369, 264288, 21906766, 21906767, 21906769, 35695917, 265021, 264018, 52278028, 246631, 265021, 264091, 65274620, 18108368, 263372, 18108376, 35696423, 224600, 22279000, 222	5264507, 52645166, 52646365, 52646842. 22278994, 56994075, 22278997, 22278998, 22278999, 60432049, 264259, 29331822, 29331824, 66714117, 29331826, 60432289, 29331827, 29331827, 29331828, 35696052, 264908, 66712502, 264909, 265008, 265009, 265009, 265009, 265009, 265009, 265009, 265009, 265009, 265009, 265009, 26640375, 264428, 264388, 2644637, 5264428, 265021, 265	UNCLASSIFIED 284288	eph 264288, 264686, 264767, 22279002
1621  87076708 (3241, 3242)	94741739 (3243, 3244) Novei Protein sim. GBank gi calmodulin-binding protein (	1623 87779106 (3245, 3246) Novel Protein sim, GBank gij731088jspjP40389jUV22_SCHPO - UV-INDUCED PROTEIN UVI22	1624 87338178 (3247, 3248) Novel Protein sim. GBank gij3875666 emb CAB05478  - (Z83104) cDNA EST EMBL:100015 comes from this gene; cDNA EST EMBL:D33665 comes from this gene; cDNA EST EMBL:D356540 comes from this gene; cDNA EST yk240f8.3 comes from this gene; cDNA EST comes from this gene; cDNA EST yk240f8.3 comes from this gene; cDNA EST yk240f8.3			83368773 (3253, 3254)	1628   85708459 (3255, 3256) Novel Protein sim. GBank gij3668087 (AC004667) - hypothetical protein [Arabidopsis thallana]

84993841 (3257, 3258)	1629 [84993841 (3257, 3258) Novel Protein sim. GBank oil4240175idbilBAA74866 11		struct	264555
87779027 (3259, 3260)			UNCLASSIFIED	29331822, 29331827, 265010, 264693, 264634, 22279002
7758454 (3261, 3262)	87758454 (3261, 3262) Novel Protein sim. GBank gil1915892 emb CAA69995  - (Y08740) tom-1A protein [Gallus gallus]		UNCLASSIFIED	55811957, 264259, 33657023, 264693, 29331822, 29331824, 29331827, 29331828, 264906, 264908, 55811576, 264910, 264634,
			-	264636, 264637, 56182323, 264559, 264758, 18108385, 264563, 264764, 264766
7871692 (3263, 3264)	87871692 (3263, 3264) Novel Protein sim. GBank gi[2558501 dbj BAA22896  - (D63850) hepatoma-derived growth factor (Mus musculus)		UNCLASSIFIED	264687, 264769, 264691, 264692, 29146499, 264509, 264905, 264907, 284511, 284512, 264807, 284511, 284512, 264897, 284511, 284897, 2848
87773683 (3265, 3266)				264488, 264859, 264907, 264908, 264909, 264628, 264629, 264831
15992817 (3267, 3268)	85992817 (3267, 3268) Novel Protein sim. GBank gi 4887229 gb AAD32244.1 AF15075 - (AF150755) microtubule-actin crossiinking factor [Mus musculus]	Contains protein domain (PF00435) - struct Spectrin repeat	struct	265007, 264637, 22279002
94232600 (3269, 3270)			UNCLASSIFIED	65274572, 22278998, 35696052, 52844045, 264511, 265008, 285009, 285010, 285011, 265018, 265019, 28448, 284369, 21906765, 21906768, 265021, 284660, 284482
80413227 (3271, 3272)			UNCLASSIFIED	22278995, 264594, 264763, 265020
0070435 (3273, 3274)	80070435 (3273, 3274) Novel Protein sim. GBank gil4557511 freflNP_001339.1 pDAPK - death-associated protein kinase 3	Contains protein domain (PF00069) - kinase Eukaryotte protein kinase domain	tinase	264558
7101854 (3275, 3276)	87101854 (3275, 3276) Novel Protein sim. GBank gil3420051 (AC004680) - unknown protein [Arabidopsis thaliana]	,		21906765, 21906767, 22278996, 35696286, 22278999, 264259, 264692, 264693, 29331824, 33657109, 264508, 264906, 18108370, 264629, 265007, 33657402, 21906754, 264602, 264604, 264764, 264683, 264566, 264288
м322194 (3277, 3278)	1639   94322194 (3277, 3278) Novel Protein sim. GBank gil5420389 emb CAB46680.1  - (AJ243460) proteophosphoglycan [Leishmania major]	·	UNCLASSIFIED	264488, 18108394, 65274572, 56182575, 35696286, 29331824, 29331826, 29331827, 35696052, 264908, 56182435, 264112, 265006, 265006, 264757, 264758, 55811386, 264603, 264760, 18108351, 264764, 264288, 264691, 33657023, 65274620, 18108370, 55810764, 55811576, 264558, 264639, 83373044, 18108385, 87168518

56182575, 56994075, 35696286, 60432049, 60432289, 29331827, 35696052, 52644045, 56182435, 264510, 265006, 265007, 285008, 265010, 265009, 33657402, 55812038, 22644229, 21906785, 21906766, 21908768, 35695017, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 265011, 26501109, 33657109, 265008, 265009, 265008, 265008, 265009, 265008, 265008, 265008, 265009, 265009, 265008, 265008, 265009, 265009, 265008, 265009, 265000, 265000, 265000, 265000, 265000,	25812038, 265010, 265011, 264555, 264556, 264558, 18108383, 25278899, 28147620, 29331826, 29331828,	33656970, 55812038, 285010, 265018, 265019, 265019, 18108351, 264689, 265020, 265022, 264690, 33657023, 65274620, 35695763, 52644332, 18108381, 60170394, 56182323, 18108388, 87168518, 222739002, 264564		264905, 264907, 264906, 264976, 205055, 264765, 264768, 264769, 264769, 264628, 264630, 264631, 264637, 264567, 264565, 264567	T			265009, 264566, 55811957, 35833917, 55810764, 284556, 56182323, 264558. 18108385
UNCLASSIFIED				UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	- UNCLASSIFIEG
	COLUMNIA (COLUMNIA COLUMNIA CO	Contains protein domain (r.r.) 1953) Suruce		-				Contains protein domain (PF00096) - UNCLASSIFIED Zinc finger, C2H2 type
94143185 (3279, 3280) Novel Protein sim. GBank gij2842469 emb CAA16847.1  - (AL021747) hypothetical protein (Schizosaccharomyces pombe]		94312557 (3283, 3284) Novel Protein sim. GBank gij1575333 (U60416) - myr 6 myosin heavy chain [Rattus norvegicus]				95362691 (3291, 3292) Novel Protein slm. GBank gij 1076802lpiri  S49915 • extensin like protein - maize	94278428 (3293, 3294) Novel Protein sim. GBank gij5002573jembjCAB44338.1  - (Y17466) alpha-N-acetylgalactosamine alpha-2.6- stalvitransferase Fuou rubrinesi	_
94143185 (3279. 3280)   N	87625160 (3281, 3282)		94131766 (3285, 3286)	88095125 (3287, 3288)				1648 87642098 (3295, 3296)
1840	<u>18</u>	1642	1643	1844	1645	1646	1647	20

	10000 10000	141	Č	- Contract	JEAABB JJJJBOOK JKROEJRK JJJJROOK
200	1050 (3531, 3530)	Sociations (Seer, Sees) Movel Protein Sint. Obsini Bijostocojaminjostocost.	<u> </u>		22278997, 22278999, 60432049, 264259,
					20111822 20111824 20111825 20131826
					20011011, 100011011, 100011010, 100011010,
					23331021, 23331020, 204303, 204301,
					66/12502, 264908, 52644045, 264909,
				-	56182435, 264511, 265007, 265008, 265009.
					264591, 264593, 60433438, 264596,
_				٠	55812038, 21906754, 265011, 284601,
		-			284602, 265017, 265018, 265019, 264682,
					264448, 264764, 264683, 264288, 284766,
					264685, 264687, 264768, 264688, 264769,
_					52644229, 264689, 21906765, 21906766,
				. •	21906767, 21906768, 55811957, 35695917,
					265021 265022 52644150 264692
					33657023 33657109 20281149 18108370.
_					20.000 10.0001 10. E0001 10. 10.0001 0.
			•		264628, 18108374, 18108378, 35696423,
_					35695855, 264632, 264634, 264635, 264636,
					18108380, 264639, 264558, 18108382,
					18108384, 18108385, 18108387, 264080,
					264404, 60432113, 22279000, 22279002,
					284482 284565 264566 264487
	COCC OUCC OCSO				285011 264602 21006767 18108374
16/8 0001	87418538 (3288, 3300)	_	,		ZODULI, ZOMOUZ, ZIBUDIOI, IOLUGAIM.
		(AL031644) possible zinc-finger protein			18108377, 18108385
1864 0162	2002 4020/ 2200	Osepate 2001 Name Desiring Charles Completed (1989) 2001 11		cychoco	254488 52545158 18108397 35595286
	39773 (3307, 3302	(A) 160037 hundhelical policy (Home emisse)		200	22278998 22278999 264259 29331822
_					20234824 20234824 20234828 20334827
					23531024, 23531025, 23531020, 23531027,
_					2331040, 204300, 204300, 43331030,
					264910, 60432229, Z1906/54, Z65010,
					265011, 265017, 265019, 264448, 18108354,
					264288, 264688, 21806765, 21906768,
_					21906768, 21906769, 265022, 264692,
					264693, 264629, 35695855, 264556, 264637,
					264557, 264559, 83373044, 56526486,
					22279000, 22279002, 264564
1652 8659	98622 (3303, 3304	86598622 (3303, 3304) Novel Protein sim. GBank gil1657837 (U73200) - p116Rip	Contains protein domain (PF00169) - struct	struct	22278997, 29146498, 56182435, 21906754,
		[Mus musculus]	PH domain		264369, 21906765, 21906768, 21906769.
					265020, 52644150, 33657109, 22279000,
					22279002
1653 9425	55993 (3305, 3306	94255993 (3305, 3306) Novel Protein sim. GBank gij3776054 jemb CAA06273] -	Contains protein domain (PF00047) - glycoprotein	glycoprotein	18108398, 22278995, 22278998, 264259,
		(AJ004999) Tapasin [Gallus gallus]	Immunoglobulin domain		29331822, 29331824, 60432289, 29331826,
					29331827, 29331830, 264909, 265006,
					265009, 60432229, 60433356, 60433438,
					21906754, 265017, 265019, 264448, 264683.
					264288, 265021, 265022, 264692, 18108364,
$\neg$					65274791, 18108384, 60432113, 264567
1654 7975	79756471 (3307, 3308)	](6		UNCLASSIFIED	33657109, 264565

5264507, 35696286, 22278998, 22278999, 29331824, 29331825, 29331828, 33656970, 264908, 52644045, 264511, 264910, 52646317, 264288, 52644229, 33657023, 33657109, 52644332, 264557, 56182323, 56526486, 60432113	29331822, 29331824, 29331825, 264563	29331627, 265009, 21906766, 21906767, 265020, 265022, 33657109, 264638, 56526486, 264482	52646365, 35696286, 22278996, 22278997, 22278998, 22278999, 264259, 29331822, 29331824, 29331825, 29331826, 29331822, 29331828, 29331828, 29331828, 29331828, 29331828, 29331828, 29331828, 2644045, 265006, 60433356, 264757, 2644348, 264369, 264288, 264766, 264768, 21906767, 21906768, 21906767, 21906768, 21906767, 21906768, 21906767, 21906768, 21906767, 21906768, 21906767, 21906768, 21906769, 21906767, 21906768, 21906769, 21906769, 21906769, 250971, 18108362, 265091, 265091, 265091, 265091, 265091, 265091, 264482	52646317, 21906766, 21906767, 21906768, 87168518, 22278996, 265020, 22278999, 87168559, 264603, 265017, 264631, 265018, 265019, 22278002, 264482, 264635, 264565	264488, 35696286, 264259, 35696052, 264508, 264905, 264906, 264906, 264907, 264908, 264906, 264906, 264907, 264908, 264906, 264907, 264908, 264511, 265008, 265007, 264501, 264501, 264501, 264501, 264501, 264501, 264501, 264501, 264501, 264501, 264501, 264601, 264601, 264601, 264691, 264691, 264691, 264691, 3695855, 264634, 264659, 264659, 264559, 264559, 264561, 2
synthase	UNCLASSIFIED	dna_ma_bind	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED
		Contains protein domain (PF00076) - Idna_ma_bind RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)	Contains protein domain (PF00829) - UNCLASSIFIED Ribosomal prokaryotic L21 protein		
1655   86689346 (3309, 3310) Novel Protein sim. GBank gi[3355717 emb CAA73496  - (Y13053) seryl-IRNA synthetase [Zea mays]	78962297 (3311, 3312) Novel Protein sim. GBank gil1890141 dbj BAA18947  - [(D83206) P24 protein [Mus musculus]	87771994 (3313, 3314) Novel Protein sim. GBank gil4557645 pef NP_001524.1 pHNRP - heterogeneous nuclear ribonucleoprotein L	87773778 (3315, 3316) Novel Protein sim. GBank gij3877072 emb CAA87060  - (Z46937) similarity with ribosomal protein L21 [Caenorhabditis elegans]	88230101 (3317, 3318) Novel Protein sim. GBank gij539218 pir  538038 - hypothetical protein YKL201c - yeast (Saccharomyces cerevisiae)	94315313 (3319, 3320) Novel Protein sim. GBank gil2497012!sp Q10010 YSV4_CAEEL - HYPOTHETICAL 26.6 KD PROTEIN T19C3.4 IN CHROMOSOME III
86689346 (3309, 3310	79962297 (3311, 3312	87771994 (3313, 3314	87773778 (3315, 3316	88230101 (3317, 3318	94315313 (3319, 332)
1655	1656	1657	1658	1659	1660

264488, 29331826, 29331828, 264509, 264906, 264909, 264909, 264510, 264911, 264910, 264511, 264910, 264593, 264595, 264758, 264596, 264760, 264760, 264766, 264768, 264629, 264630, 264667, 264686, 83373044, 264564, 264566, 264567, 264486	56994075, 22278999, 264259, 29331824, 29331826, 29331827, 29146498, 265009, 33109954, 87168559, 265019, 264288, 264688, 21806767, 21806769, 264691, 33857182, 18108370, 18108374, 18108385, 22279002	264259, 29331822, 29331826, 264905, 264906, 264908, 264908, 264510, 265001, 265001, 265016, 265011, 87168559, 265017, 265018, 265019, 26448, 264766, 264686, 21906765, 21906767, 21906769, 265020, 265021, 26170615, 264690, 264692, 264693, 264655, 83373044, 60432113, 22279000, 22279002	66714117, 264508, 264509, 264906, 264907, 264908, 264907, 264908, 264511, 264910, 264764, 264687, 264689, 33657109, 35696423, 35695855, 264632	29331825, 33109954, 264369, 264767, . 264669, 33657109, 83373044	264259, 29331824, 29331827, 60433438, 265022, 264638	35696286, 22278997, 29331825, 264909, 21906754, 265017, 265018, 265019, 264682, 264683, 264768, 264688, 21906766, 21906767, 21906768, 21906769, 264691, 264555, 264556, 22279000, 264566	264906, 264909, 264632, 18108381
immunoglob	helicase	transport	kinase	UNCLASSIFIED	UNCLASSIFIED	kinase	deaminase
Contains protein domain (PF00047) - immunoglob Immunoglobulin domain	Contains protein domain (PF00575) - helicase S1 RNA binding domain	Contains protein domain (PF01412) - transport Putative GTP-ase activating protein for Arf				Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	
1665   91226952 (3329, 3330) Novel Protein sim. GBank gil 1083506 pir  S50065 - slaloadhesin - mouse	95358160 (3331, 3332) Novel Protein sim. GBank gij3913431[sp]042643[DDX8_SCHPO - PUTATIVE PRE- MRNA SPLICING FACTOR ATP-DEPENDENT RNA HELICASE SPAC10F6.02C	91228655 (3333, 3334) Novel Protein sim. GBank gil5689535 dbj BAA83051.1  - (AB029022) KIAA1099 protein [Homo sapiens]	88095135 (3335, 3336) Novel Protein sim. GBank gij2076894[gb]AAB53983.1] - (AF002197) short region of weak similarity to protein kinase C; contains similarity to Pfam domain PF00130 (DAG_PE-bind). Score=10.0. E-value=0.0034, N=1 [Caenorhabditis elegans]	91227846 (3337, 3338) Novel Protein sim. GBank gij3875371[emb]CAA85414.1[		87346372 (3341, 3342) Novel Protein sim. GBank gil462451 spl934244 KKK1_YEAST - PROBABLE SERINE/THREONINE-PROTEIN KINASE YKL101W	88291834 (3343, 3344) Novel Protein sim. GBank gij1814270 (U74586) - double- stranded RNA specific adenosine deaminase [Rattus
91226952 (3328, 3330) h	95358160 (3331, 3332) I	91228655 (3333, 3334)	88095135 (3335, 3336)	91227846 (3337, 3338)	87628009 (3339, 3340)	87346372 (3341, 3342)	86291834 (3343, 3344)
1665	1668	1667	1668	1669	1670	1671	1672

1673	88095137 (3345, 3346)	1673 [88095137 (3345, 3346] Novel Protein sim. GBank gi 2076894 gb AAB53983.1  - (AF002197) short region of weak similarity to protein kinase C; contains similarity to Pfam domain PF00130 (DAG_PE-bind), Score=10.0, E-value=0.0034, N=1 [Caenorhabditis elegans]	Contains protein domain (PF00130) - kinase Phorbol esters/diacy/glycerol binding domain (C1 domain)	kinase	264488, 264569, 18108394, 56934075, 22278996, 264559, 35696052, 264508, 264908, 264905, 264907, 264909, 264909, 264510, 264910, 60170831, 264592, 264595, 264693, 264693, 264683, 2646
					264686, 264768, 264687, 264769, 264689, 264690, 33657023, 264692, 264693, 33657109, 264628, 264629, 18108374, 35696423, 35695855, 264631, 264632, 264634, 264635, 264563, 264563, 264563, 264566, 264567, 264486, 264567, 264486
1674	88258028 (3347, 3348)	88258028 (3347, 3348) Novel Protein sim. GBank gil5262467 emb CAB45693.1  - (AL080062) hypothetical protein [Homo sapiens]		kinase	29331822, 29331824, 264906, 52644045, 60433356, 87168559, 264448, 264288, 264686, 264691
1675	87606466 (3349, 3350)	87606466 (3349, 3350) Novel Protein sim. GBank gij3128366 (AF010496) - 50S ribosomal protein 19 [Rhodobacter capsulatus]		UNCLASSIFIED	56181686, 35696286, 22278997, 22278998, 264259, 29331824, 29331827, 35696052, 66712502, 264764, 264288, 264686, 264687, 35695917, 255020, 264690, 284693, 35695763, 18108370, 35696423, 3569585, 264639, 18108385, 264564
1676		95358086 (3351, 3352) Novel Protein sim. GBank gil4164065 gb AAD05327  - (AF111091) latrophilin 3 splice variant bbaf [Bos taurus]		UNCLASSIFIED	264259, 29331827, 29331828, 264106, 264907, 265009, 264000, 265019, 264288, 21906765, 265020, 265022, 35695855, 83373044, 18108385
1677	87408587 (3353, 3354)	87408587 (3353, 3354) Novel Protein sim. GBank gil3327046 dbj BAA31591  -		UNCLASSIFIED	264908
10/8	46866829 (3355, 3356)			UNCLASSIFIED	29331824, 264102
	91214106 (3357, 3358)	91214108 (3357, 3358) Novel Protein sim. GBank gil550452 (U08469) - 3- methylcrotonyl-CoA carboxylase, biotin-carrier domain [Glycine max]	Contains protein domain (PF00289) - carboxylase Carbamoyl-phosphate synthase (CPSase)	carboxylase	264468, 16108392, 16108394, 52646842, 18108397, 18108398, 35696286, 29331824, 265006, 265007, 265008, 265009, 18108348, 265011, 18108351, 264683, 18108354, 18108356, 18108359, 21906765, 29148627, 29148629, 264690, 18108361, 18108362, 18108364, 18108365, 18108368, 264628, 18108379, 35696423, 35695855, 264638, 18108385, 18108382, 18108384,
1680		91005372 (3359, 3360) Novel Protein sim. GBank gi[2394478 (AF024500) - No definition line found [Caenorhabditis elegans]		transport	65274572, 22278994, 22278999, 66714117, 29331827, 56182435, 21906754, 265018, 264288, 21906769
1681		94324150 (3361, 3362) Novel Protein sim. GBank gij5689537 dbjjBAA83052.1  - (AB029023) KIAA1100 protein [Homo saplens]		UNCLASSIFIED	22278996, 29331822, 264908, 264593, 264604, 265019, 264608, 55811957, 264690, 31657023, 31657023, 31657023
1682	86042710 (3363, 3364)			UNCLASSIFIED	264909, 265017, 284605

263994, 35696286, 35696052, 264508. 264509, 264905, 264906, 264907, 264908. 264909, 264511, 265006, 265007, 265009, 264910, 264593, 264758, 265010, 265011, 264760, 264761, 264762, 264763, 284764, 264369, 284766, 264768, 35695917, 264692, 33657109, 35696423, 35695855, 264634, 264635, 264636, 264638, 264639, 83373044, 264486	264563, 264566	60424179, 52646842, 22278994, 35696286, 22278998, 264259, 52645080, 29331824, 29331826, 265007, 33657084, 265018, 264681, 264448, 264683, 264369, 21906765, 21906768, 21906769, 265021, 264692, 65274620, 33657109, 2749626, 2644332, 56182323, 22279000	56182575, 264259, 264508, 264905, 264509, 264907, 264908, 264510, 264511, 265006, 264512, 265009, 264510, 264758, 265010, 265011, 264605, 18108351, 264764, 264766, 18108357, 264628, 264630, 264631, 264634, 264635, 264639, 264639, 264636, 264565, 264486, 264567	35695917, 264906, 264907, 264908, 264510,	265006, 265007, 264910, 264556, 18108381, 18108383, 265011	264905, 264910, 264760, 264629, 264555	264569, 35696286, 22278999, 264508, 264908, 264909, 56182435, 33657402,	6043,3436, 55612,038, 2550117, 2550136, 266448, 264764, 264288, 264688, 29148629, 35695917, 265020, 265021, 263972, 18108374, 65274791, 83373044, 264089	264369, 21906766, 264692, 264639,	87168518	264768, 33657109, 29331827, 29146629. 264510, 264106, 264910, 264109, 264508. 60170831, 264563, 264905, 264564, 264691.	264637, 264628, 264907, 264908, 33657023, 284567, 264768, 263974	264106	29331826, 264508, 264905, 264907, 264595, 265010, 265011, 21906768, 33657023, 264629, 263978, 264558
UNCLASSIFIED	UNCLASSIFIED	kinase	UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED		cadherin					UNCLASSIFIED
Contains protein domain (PFD1852) - UNCLASSIFIED START domain		Contains protein domain (PF00625) - kinase Guanylate kinase							Contains protein domain (PF00028) - cadherin	Cadherin domain				Contains protein domain (PF00042) - UNCLASSIFIED Globin
1683   94316213 (3365, 3366)   Novel Protein sim. GBank gij5031717 ref NP_005704.1 pGPBP - goodpasture antigen-START domain binding protein		Novel Protein sim. GBank gi 1255371 (U53147) - coded for by C. elegans cDNA yk34a9.5; coded for by C. elegans cDNA yk34a9.3; Similar to guanylate kinase. [Caenorhabditis elegans]	1686   87820710 (3371, 3372)   Novel Protein sim. GBank gi[2244707 dbj  BAA21115.1  - (AB005287) thrombospondin 1 [Bos taurus]	94719400 (3373, 3374) Novel Protein sim. GBank	gi 4680679 gb AAD27729.1 AF13295 - (AF132954) CGI-20 protein [Homo sapiens]		94325049 (3377, 3378) Novel Protein sim. GBank gil4240193 dbj BAA74875.1  - (AB020659) KIAA0852 protein [Homo sapiens]		83255346 (3379, 3380) Novel Prolein sim. GBank gij3800736 (AF031572) - seven-	pass transmembrane receptor precursor [Mus musculus]	88095223 (3381, 3382) Novel Protein sim. GBank gij2773208 (AF039713) - No definition line found (Caenorhabditis elegans)		10	87012775 (3385, 3388) Novel Protein sim. GBank gi 121271 sp P02207 GLB_LAMFL · GLOBIN
94316213 (3365, 3366)	80063409 (3367, 3368)		87820710 (3371, 3372)	94719400 (3373, 3374)		82158442 (3375, 3376)	94325049 (3377, 3378)		83255346 (3379, 3380)		88095223 (3381, 3382)		86106709 (3383, 3384)	87012775 (3385, 3386)
5 88	1684 E		989	1687		_	1689		1690		1691		1692	

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35696286, 22278998, 22278998, 22278999, 264259, 29331822, 29331825, 29331826, 28331826, 28331826, 28331826, 28331826, 28331828, 28331828, 28331828, 28331828, 284390, 284490, 284592, 60433356, 21906754, 265010, 265011, 18108351, 264768, 264689, 21906768, 60170615, 264691, 264692, 36696423, 35695855, 264556, 18108381, 18108356, 87166518, 264486	264634	22278997, 264259, 265010, 18108351, 264764, 21906766, 18108370	264682	56994075, 22278998, 22278999, 264509, 33657402, 264758, 87168474, 87168559, 265017, 265018, 26448, 264687, 29148627, 21906769, 29148629, 265020, 265022, 33657023, 264558, 87168518, 22279002	35696286, 264635	29331824, 52644045, 265008, 265009, 263969, 263971	264092, 264110, 263977	22278995, 22278997, 264092, 29146498, 29146499, 264107, 264508, 264907, 264110, 2641112, 265009, 60170831, 21906754, 265011, 265017, 264762, 18108351, 264288, 21906765, 35695917, 265021, 60170615, 263967, 33657109, 18108370, 263972, 263974, 18108374, 283976, 356526486, 87168518, 60432113	264909, 265017, 264628, 264629, 264638	264369	264569, 35696286, 264907, 265010, 264687, 264768, 264692, 264693, 264636, 264566	22278996, 22278998, 264259, 264509, 265018, 265018, 264685, 264686, 21906768, 21906769, 265022, 264691, 264558, 22279000
mapolymerase	UNCLASSIFIED		glycoprotein		UNCLASSIFIED	nucl_recpt	UNCLASSIFIED	МНС	UNCLASSIFIED	UNCLASSIFIED		н
Contains protein domain (PF01193) - mapolymerase RNA polymerases L / 13 to 16 kDa subunit				,		Contains protein domain (PF00096) - nucl_recpt Zinc finger, C2H2 type						
5.1[pPOLR - polymerase (RNA) II e J (13.3kD)	94719325 (3389, 3390) Novel Protein sim. GBank gi 4680679 gb AAD27729.1 AF13295 - (AF132954) CGI-20 protein [Homo saplens]	87824038 (3391, 3392) Novel Protein sim. GBank gil4220517 emb CAA22990  - (AL035356) hypothetical protein [Arabidopsis thaliana]	85740963 (3393, 3394) Novel Protein sim. GBank gij505652 (U10362) - GP36b glycoprotein [Homo sapiens]	87445285 (3395, 3396) Novel Protein sim. GBank gi 5052031 gb AAD38411.1 AF15573 - (AF155739) axotrophin [Mus musculus]		87859161 (3399, 3400) Novel Protein sim. GBank gij543344 pir  S41647 - zinc finger 5 protein - mouse		87795092 (3403, 3404) Novel Protein sim. GBank gij3877439 emb CAA96652  - (Z72510) similarity to yeast UTR3 protein (Swiss Prot accession number P21374); cDNA EST EMBL:D72822 comes from this gene; cDNA EST EMBL:D75763 comes from this gene; cDNA EST yk274e3.3 comes from this gene; cDNA EST yk274e3.3 comes from this	79568651 (3405, 3406) Novel Protein sim. GBank gil451544 (U04267) - proline-rich  cell wall protein [Gossyplum barbadense]	86622979 (3407, 3408) Novel Protein sim. GBank gi[1263289 (U47856) - fibroin-4 [Araneus diadematus]	87795175 (3409, 3410) Novel Protein sim. GBank gil4519621 db  BAA75670.1 -   (AB017614) OASIS protein [Mus musculus]	1706   87790967 (3411, 3412) Novel Protein sim. GBank gij3123034 sp Q15011 Y025_HUMAN - HYPOTHETICAL. PROTEIN KIAA0025
1694   94208168 (3387, 3388) Novel Protein sim. GBank gij6453932 reflNP_00622( (DNA directed) polypeptid	94719325 (3389, 3390)		85740963 (3393, 3394) I	87445285 (3395, 3396)			86570488 (3401, 3402)					87790967 (3411, 3412)
1694	1695	1696	1697	1698	1699	1700	1701	1702	1703	1704	1705	1708

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1717	87563223 (3433, 3434)	1717   87553223 (3433, 3434)   Novel Protein sim. GBank gi[2755411[emb]CAA74749  -		UNCLASSIFIED	264569, 264259, 29331825, 29331826, 29331828, 35696052, 264509, 264905
					264907, 264908, 264909, 264512, 265009,
					264910, 264592, 264595, 264758, 264759,
					265017, 264681, 264764, 264766, 264686,
					18108357, 35695917, 264690, 264692,
					264693, 264628, 264629, 35696423, 264630,
					264631, 264635, 264636, 18108380, 284638,
1	ı				264639, 18108388, 18108391
1718		87032628 (3435, 3436) Novel Protein sim. GBank		UNCLASSIFIED	265011, 264681, 264682, 264684, 264688,
		gil2833262 sp Q14999 Y076_HUMAN - HYPOTHETICAL			264689, 21906765, 265021, 264691,
		PROTEIN KIAA0076 (HA0936)			33657023, 264693, 18108370, 35695855.
_			.:		264632, 264634, 264636, 18108388,
					22279002
1719	Ī	94315259 (3437, 3438) Novel Protein sim. GBank	,	UNCLASSIFIED	18108396, 65274572, 35696286, 22278997,
		gi[4505197]ref[NP_003473.1]pMLL2 - myeloid/lymphoid or			60432049, 56182181, 66714117, 60432289.
		mlxed-lineage leukemia 2			29331826, 35696052, 29331828, 264906,
					29331830, 56182435, 264592, 60431735,
					60433438, 55812038, 264759, 265010,
					264600, 264601, 265017, 264448, 264764,
					284288, 264769, 21906766, 21906769,
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					33657023, 33657109, 33657182, 27486262,
					33657349, 35695763, 18108370, 60431528.
					18108374, 35696423, 55811576, 35695855,
					264631, 56182323, 264559, 264564, 264486
1720	_	94853063 (3439, 3440) Novel Protein sim. GBank gij2129478 pir  S51939 -		UNCLASSIFIED	56182575, 22278999, 264259, 29331824,
		chilinase (EC 3.2.1.14) precursor - beet			60432289, 29331827, 35696052, 264508,
					264905, 264906, 264907, 264908, 264909,
					264511, 264910, 264758, 21906754, 265011.
_					264601, 264760, 264762, 264288, 264766.
	-		_		264686, 18108357, 264689, 21906765,
					55811957, 264693, 20281149, 264629,
					18108374, 55811576, 65274791, 264630,
					20281071, 264634, 264635, 264636, 264637,
					264556, 264638, 264639, 56182323,
	_				87168518
1721				UNCLASSIFIED	22278994, 22278999, 29331822, 265006,
<u>.</u>		(AL050280) hypothetical protein [Homo sapiens]			265007, 265008, 55812038, 21906754.
					60174639, 265011, 87168559, 18108351,
		*			18108354, 21906765, 21906766, 21906768,
					21906769, 265020, 33657109, 18108370,
				•	18108374, 264556, 60170394, 83373044,
					18108385, 264486
1722		94134549 (3443, 3444) Novel Protein sim. GBank gij5689375 dbj BAA82968.1  -	Contains protein domain (PF00567) -	kinase	56994075, 29331824, 29331828, 265009,
		(AB030644) tudor repeat associator with PCTAIRE 2	Tudor domain		18108351, 21906768, 265020, 33657023.
		[Rattus norvegicus]			18108374, 83373044

97. 25. 49. 007. 592. 84.	631.	77. 11. 1764. 18.		
264488, 264687, 264769, 21906767, 21906768, 56182575, 55811957, 22278997, 22278997, 22278998, 265020, 264259, 264692, 33657023, 29331822, 29331824, 29331825, 60432289, 33657182, 33856970, 3365739, 2946499, 264508, 264907, 18108374, 55811576, 264510, 265001, 264591, 264591, 264591, 264591, 264591, 264591, 264591, 265001, 265007, 264591, 264591, 265001, 265001, 265001, 265019, 264563, 264764, 26488, 264369, 264766	264905, 18108359, 264693, 264628, 264631, 264638, 264555, 264556, 264558, 264559	35696286, 264259, 29331822, 35696052, 264508, 264509, 264905, 264905, 264906, 265009, 264907, 264901, 265009, 264901, 264001,	29146498, 264683, 264689	264905, 265011, 264689, 21906768
UNCLASSIFIED			UNCLASSIFIED	UNCLASSIFIED
	gi 2340162 (AF005083) - dsRBP- Contains protein domain (PF00096) - UNCLASSIFIED Zinc finger, C2H2 type	Contains protein domain (PF00023) - homeobox Ank repeat	•	
1723 95358181 (3445, 3446) Novel Protein sim. GBank gil4426962 gb AAD20633  - (AF126062) Arf-like 2 binding protein BART1 [Homo sapiens]		Novel Protein sim. GBank gij3152662 (AF064604) - KE03 protein [Homo sapiens]		85296362 (3453, 3454) Novel Protein slm. GBank gil4689348lgblAAD27861.1lAF13256 - (AF132562) BCDNA.LD14270 [Drosophila melanogaster]
95358181 (3445, 3448) [	87713806 (3447, 3448)	85655191 (3449, 3450)	85754255 (3451, 3452)	, 85296362 (3453, 3454)
1723 (	1724	1725	1726	

9515 (3455, 3456)	1728  95349515 (3455, 3456) Novel Protein sim. GBank gil4406549 gb AAD20027 -	i i	UNCLASSIFIED	60424179, 18108397, 56182575, 22278995, 56994075, 35696286, 22278997, 22278998,
	(AF13175b) Unknown (Trong Saprens)			22278999, 264094, 60432049, 264259. 29331822, 29331824, 56182181, 29331825, 60432289, 29331824, 56182181, 29331825, 60432289, 29331826, 29331827, 35696652, 264095, 264907, 29331830, 66712502, 264908, 264907, 29331830, 66712502, 265009, 6043345, 264511, 265017, 265019, 264763, 264683, 264681, 264684, 264369, 264685, 264763, 264683, 264687, 264684, 264369, 21906764, 264689, 21906764, 264689, 21906764, 264689, 21906764, 264639, 264692, 33657023, 264692, 33657023, 264693, 33657109, 18108370, 264638, 264634, 264634, 264634, 264634, 264634, 264634, 264634, 264634, 264634, 264638, 83373044, 56526486, 87168518, 60432113, 22279000, 22279002, 264564, 264566
7, 3458)	91227948 (3457, 3458) Novel Protein sim. GBank gil854065[emb CAA58337] - (789413) HRR Pluman hemesylus 6]		UNCLASSIFIED	264906, 264907, 264908, 264511, 264555. 83373044, 264596, 264566
85483474 (3459, 3460)			UNCLASSIFIED	29331822, 29331825, 29331828, 264907, 264908, 264909, 265011, 264764, 264629
31, 3462)	88266068 (3461, 3462) Novel Protein sim. GBank gij631600 pir  S47094 - hypothetical protein - rabbit	,	UNCLASSIFIED	52646842, 264907, 284909, 56182435, 55811386, 87168559, 285018, 265019, 264760, 52644229, 55811576
63, 3464)	91218878 (3463, 3464) Novel Protein sim. GBank gil4240231 db  BAA74894.1  - (AB020678) KIAA0871 protein (Homo saplens)		struct	56182575, 29331822, 29331824, 29331824, 66712502, 264591, 33657402, 60433356, 265019, 21906768, 21906769, 35695917, 265020, 265021, 264638, 56182323
165, 3468)	87617178 (3465, 3466) Novel Protein sim. GBank gij1575756 (U70674) - m-Numb [Mus musculus]	Contains protein domain (PF00640) - synthase Phosphotyrosine interaction domain (PTB/PID).	synthase	264807, 264910, 33657402, 265010, 264001, 264663, 264684, 264686, 264769, 264691, 264692, 264693, 264628, 264536, 284556
87795261 (3467, 3468)				284693

264488, 18108394, 18108398, 22278998, 60432049, 264259, 29331822, 264908, 265006, 265007, 265008, 265009, 264591, 33657402, 285010, 265011, 87168559, 265017, 18108351, 264682, 18108354, 264692, 2846829, 21906768, 21906768, 21906768, 265021, 265022, 52644150, 18108364, 18108386, 18108350, 18108350, 18108359, 18108359, 264480, 264259, 66714117, 66712502,	58182435, 265008, 265008, 264910, 60433356, 87168559, 265017, 285019, 18108351, 265020, 265022, 33657023, 33657109, 264555, 60431850, 264637, 60170394, 264558, 264639, 264634, 264564	265018, 265019, 22279002	264908, 264909, 265008, 264910, 264300	35696052, 264603, 264557	264604, 21906764, 18108364, 264629. 35695855, 264636	60432289, 29331827, 264509, 265009, 60432229, 264759, 265017, 264767, 264688, 264689, 21906769, 265020, 33657109	264908, 264910, 264758, 265011, 264631, 264638, 264566	265017, 265020, 264692	22278998, 264508, 264907	264558	65274791, 264639, 264559	52646842, 29331824, 29331825, 52644045, 56182435, 265007, 52646317, 87168474, 265018, 18108351, 264369, 264769, 264689, 35695917, 60170615, 264691, 33657023, 65274791, 35695855, 60170394, 22279000, 264482
INCI ASSIFIED		kinase	UNCLASSIFIED	UNCLASSIFIED	traffic	UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	transcriptfactor
Contains protein domain (PF0169) - Uncharacterized protein family UPF0016		Contains protein domain (PF00168) - kinase C2 domain	200	Contains protein domain (PF00441) - UNCLASSIFIED Acyl-CoA dehydrogenase		Contains protein domain (PF01363) - UNCLASSIFIED FYVE zinc finger			*		Contains protein domain (PF00400) - UNCLASSIFIED WD domain, G-beta repeat	
AF14679 - (AF146793) PFT27			in sim. GBank gi[2225941[emb]CAA69714] - des protein [Mus musculus]		83592939 (3479, 3480) Novel Protein slm. GBank gil4809[emb CAA44309] - (X62452) YCR601 (Saccharomyces cerevisiae)	95010100 (3481, 3482) Novel Protein sim. GBank gil4883898 gbJAAD31695.1 AF13042 - (AF130420) serine protease-like protein isoform Homo saplens	85788814 (3483, 3484) Novel Protein sim. GBank gil4505193[ref]NP_003667.1[pMLD] - membrane fatty acid ffinial desaturase	4-	91224003 (3487, 3488) Novel Protein sim. GBank gi 728832 sp P39189 ALU2_HUMAN - !!!! ALU SUBFAMILY SB WARNING ENTRY !!!!		84326110 (3491, 3492) Novel Protein sim. GBank gil731756 sp P38873 YHY6_YEAST - HYPOTHETICAL stray of the protein in CND1.ix11 INTERGENIC REGION	94324333 (3493, 3494) Novel Protein sim. GBank gi 1658503 (U75467) - Atu [Drosophila melanogaster]
88318638 (3469, 3470) P	95362884 (3471, 3472)	88165549 (3473, 3474)	85788811 (3475, 3476)	87328576 (3477, 3478)	83592939 (3479, 3480)		85788814 (3483, 3484)	86966475 (3485, 3486)	+			
1735	1736	1737	1738	1739	1740	1741	1742	1743	1744	1745	1746	1747

748	88003580 (3495, 3496)	1748   88003580 (3495, 3496)   Novel Protein sim. GBank gl 4504511 ref NP_001530.1 pHSJ2 - heat shock protein, DNAJ-like 2	Contains protein domain (PF00684) - eph DnaJ central domain (4 repeats)		264489, 56182575, 29331824, 56182435, 264112, 265007, 265019, 264764, 21906768, 265020, 264691, 55811576, 264635, 264555, 264556, 264557, 264559
1749		83363091 (3497, 3498) Novel Protein sim. GBank gi 5650780 gb AAD45948.1 AF15196 - (AF151968) RGS protein RGS-17 [Gallus gallus]	Contains protein domain (PF00615) - oncogene Regulator of G protein signaling domain		264106
1750		Novel Protein sim. GBank gl 4996894 gb AAC28444.2  - (AF065164) hyperpolarization-activated, cyclic nucleotide-gated channel 2 [Homo sapiens]			33657402, 264288, 52644150, 263974, 83373044
1751		83373058 (3501, 3502) Novel Protein sim. GBank gilz760161 dbj BAA24184  - (AB010054) outer arm dynein light chain 2 [Anthoddaris crassispina]	Contains protein domain (PF00560) - ATPase_associated 265010, 264369 Leucine Rich Repeat	ATPase_associated	265010, 264369
1752		86456530 (3503, 3504) Novel Protein sim. GBank gi 3915482 sp P74346 YG29_SYNY3 - HYPOTHETICAL 36.0 KD PROTEIN SLR1629	Contains protein domain (PF00849) - deaminase RNA pseudouridylate synthase		264510, 264593, 264682, 21906765, 18108370
1753		94235159 (3505, 3506) Novel Protein sim. GBank gi 2852636 (AF007155) - unknown {Homo sapiens}	Contains protein domain (PF01553) - phosphatase Acyltransferase	phosphalase	56994075, 22278996, 264908, 60170831, 264682, 264764, 264369, 264288, 264685, 264687, 21906766, 264692, 264693, 65274620, 65274791, 35695855, 264637, 264564
1754		88095323 (3507, 3508) Novel Protein sim. GBank gil731421 sp P39981 YEH4_YEAST - HYPOTHETICAL 53.3 KD PROTEIN IN HXT8-CAN1 INTERGENIC REGION	`	transport	264488, 35696286, 264509, 264906, 264907, 264907, 264909, 264511, 264910, 264591, 33657402, 264509, 264757, 264758, 264600, 264604, 264762, 264763, 264685, 264768, 264691, 264634, 264588, 264632, 264634, 264564, 264565, 264566, 264567
1755		79470262 (3509, 3510) Novel Protein sim. GBank gi]1176422 (U43194) - rhophilin [Mus musculus]		UNCLASSIFIED	264686
1756		92962614 (3511, 3512) Novel Protein sim. GBank gil4432860 gb AAD20708  - (AC006300) putative glucose-induced repressor protein [Arabidopsis thaliana]			52646842, 22278994, 22278995, 56994075, 22278996, 22278997, 264259, 29331822, 60432289, 29331822, 60432289, 29331822, 255009, 60432229, 60433356, 60433438, 33109954, 21906754, 265017, 265018, 265019, 26448, 264369, 265020, 265021, 265022, 264692, 27486262, 27486264, 18108376, 20281152, 264558, 22730002, 26482
1757		95357380 (3513, 3514) Novel Protein sim. GBank gij5441615 emb CAB46856.1 - (AJ388557) zinc finger protein [Canis familiaris]	Contains protein domain (PF00096) - dna_ma_bind Zinc finger, C2H2 type	dna_ma_bind	22278997, 264259, 60432289, 29331827, 264908, 52644045, 265008, 264593, 265019, 264766, 21906768, 65274620, 18108385, 60432113, 264568, 264487

22278994, 22278999, 29331822, 29331824, 29331825, 33656970, 264508, 265006, 265007, 265009, 264591, 33657402, 33109954, 87168474, 264600, 265017, 265018, 21908749, 265020, 265017, 265018, 21908749, 265020, 265017, 26502	2857023, 33657109, 264629, 18108374, 35695855, 284632, 52644332, 22278002, 284563	264759	56182575, 60432049, 35696052, 264805, 264906, 264906, 264907, 264907, 264909, 265006, 265000, 264901, 60432239, 264592, 264595, 264762, 18108351, 264764, 264766, 264768, 264769, 21806765, 25811957, 35695917, 224690, 264692, 264639, 264639, 264634, 264636, 264557, 264639,	60432113, 264565, 264486	29331822, 264910, 264665, 264686	22278999, 264259, 264905, 264907, 60170831, 265010, 265011, 265017, 26448, 21906765, 21906767, 21906768, 265021, 264690, 33657109, 18108374, 264558, 60170394	56181686, 29331825, 35696052, 264905, 264906, 264906, 264909, 264763, 264682, 264769, 35695917, 265022, 33657023, 18108374, 35696423, 264634	22278998, 264259, 35696052, 29331828, 56182435, 265008, 265017, 265018, 264448, 264288, 21906766, 21906767, 29146627, 35695917, 264691, 33657023, 60432113, 22279002	264488 18108394, 22278995, 22278997, 22278899, 264259, 264104, 264508, 264905, 264907, 265008, 264910, 265002, 265010, 265010, 265010, 265010, 265010, 265010, 265010, 264903, 265018, 264288, 264766, 18108359, 21906766, 21906767, 29148627, 29148629, 35695917, 265020, 265021, 264692, 264628, 264629, 18108374, 263976, 294639, 264629, 18108374, 263976, 246636, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 264629, 262799000, 222799002
UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	desaturase			UNGLASSIFIED
					Contains protein domain (PF00618) - UNCLASSIFIED Guanine nucleotide exchange factor for Ras-like GTPases; N-terminal motif	Contains protein domain (PF00173) - desaturase Heme-binding domain in cytochrome b5 and oxidoreductases		Contains protein domain (PF01410) - Fibrillar collagen C-terminal domain	
1758   87612971 (3515, 3516) Novel Protein sim. GBank gij3881040jembjCAA16403j - (AL021497) predicted using Genefinder (Caenorhabditis elegans)		3518)	87329716 (3519, 3520) Novel Protein sim. GBank gil5262748 emb CAB45688.1  - (AJ133120) Proline rich synapse associated protein 2 (Rattus norvegicus)		87409586 (3521, 3522) Novel Protein sim. GBank gij127749 sp P10569 MYSC_ACACA - MYOSIN IC HEAVY CHAIN	95319887 (3523, 3524) Novel Protein sim. GBank gij3169158 (AC004770) - BC289730_2 [Homo sapiens]	9j224013 (3525, 3526) Novel Protein sim. GBank gil4809026jgb[AAD30062.1] - (AF132856) suppressor of G2 allele of skp1 homolog [Homo sapiens]	87757697 (3527, 3528) Novel Protein sim. GBank gij1360669jpir  CGHU1V - collagen atpha 1(V) chain precursor - human	91230091 (3529, 3530) Novel Protein sim. GBank gil486806[piri]S35503 - finger protein neuralized - fruit fly (Drosophila melanogaster)
1758   87612971 (3515,		1760 36004372 (3517 3518)			1761 87409586 (3521,	1762 95319887 (3523,	1763 91224013 (3525	1764 87757697 (3527,	1765 91230091 (3529

1766   95091201 (3531, 3532)   Novel Protein sim. GBank     1769   97755998 (3533, 3534)   Novel Protein sim. GBank     1771   94233542 (3541, 3542)   Novel Protein sim. GBank     1772   9541116824 (3545, 3546)   Novel Protein sim. GBank     1773   94116824 (3545, 3546)   Novel Protein sim. GBank     1774   94233542 (3541, 3542)   Novel Protein sim. GBank     1777   94116824 (3545, 3546)   Novel Protein sim. GBank     1777   94116824 (3545, 3546)   Novel Protein sim. GBank     1777   94116824 (3545, 3546)   Novel Protein sim. GBank gil3978464 (AF085683) - Gontains protein domain (PF01512) - Iransferase gil3914 (91998442)     1777   94116824 (3545, 3546)   Novel Protein sim. GBank gil3978464 (AF085683) - Gontains protein domain (PF01612) - Potassium_channel protein domain Pr	52645156, 87168559, 60170615, 53637023, 201602 33657400 37486264 364645	264693, 33657109, 27486261, 264555.	83373044	265017, 265019, 264686, 264768, 265020. 264692	20111824 20111825 264591 56182323	254461	373000000000000000000000000000000000000	264488, 264768, 264769, 36182373.	55811957, 264690, 264691, 35696052, 264905, 264509, 264906, 264907, 264628.	264908, 264909, 264910, 264634, 264635,	264616 264558 264757 284758 55812038.	65274444, 264760, 264563, 264762, 264764.	264684, 264766	264758, 264600, 264369, 55811957, 265020,	83373044, 22279000			22278008 28231828 33108954 265018	2227 0990, 49351020, 55103557, 400010;	265019, 264764, 21906765, 265020, 265021.	$\neg$ r		222/8999, 204239, 2333/024, 00424209.	29331623, 60432289, 33090032, 237100, 1284908, 1284908, 1284908, 1284908, 128480808, 1284808, 1284808, 1284808, 1284808, 1284808, 1284808, 128480808, 1284808, 1284808, 1284808, 1284808, 1284808, 1284808, 128480	1204309, 204300, 204300; 20201000; 200000 142644045, 264511, 265008, 265007, 265008,	60170831, 60433438, 264758, 55811388,	87168559, 265017, 264604, 265019,	55811150, 264288, 56181562, 264689.	21906766, 21906767, 21906768, 21906769,	55811957, 265020, 265022, 52644150,	264691, 33657023, 264692, 264693,	60431528, 35696423, 35695855, 264636,	56182323, 18108387, 56526486, 22279000.	22279002, 264563, 264564, 264565, 264565.	264567	65274572, 56182575, 35698052, 35812030, 33100954, 24008754, 265017, 21906767,	21906768, 21906769, 265020, 264691,	264636, 56182323, 22279002
GG_DROME - UDP- EIN GLUCOSYLTRANSFERASE gij4176443jembjCAA18263.11- (novel protein) [Homo sapiens] TAT- RAT - UDP-NPEPTIDE N- L-PEPTIDE N- L-TRANSFERASE 110 KD RANSFERASE P110 SUBUNIT) K A 1.1/AF12136 - (AF121360) nger protein 11 [Drosophila k gij3978464 (AF085693) - G kinase-associated ADP se-activating protein [Rattus	hycoprotein			JNCLASSIFIED	A COLUMN	NCLASSIFIED	JACKSSILIED	complement						ransferase								potassium_channel														UNCLASSIFIED		
GG_DROME - UDP- EIN GLUCOSYLTRANSFERASE gij4176443jembjCAA18263.11- (novel protein) [Homo sapiens] THEAT - UDP-NPEPTIDE N- LTRANSFERASE 110 KD RANSFERASE P110 SUBUNIT) K I.1JAF12136 - (AF121360) nger protein 11 [Drosophila k gij3978464 (AF085693) - G kinase-associated ADP se-activating protein [Rattus	6							Contains protein domain (PF00089) - c	Trypsin					Contains protein domain (PF00515) -	TPR Domain				Contains protein domain (PF01529) -	DHHC zinc finger domain		Contains protein domain (PF01412) -	Putative GTP-ase activating protein	for Arf					,	***						Contains protein domain (PF00415) - UNCLASSIFIED	Regulator or chromosome condensation (RCC1)	
11, 3532) 33, 3534) 36, 3540) 39, 3540) 41, 3542) 42, 3546		Ċ	gijz489067 jspjagosząjogog_brome - obi -	Novel Protein sim. GBank gij4176443 emb CAA18263.1 -	(AL022238) dJ1042K10.4 (novel protein) [Homo sapiens]			Novel Protein sim GBank	HUMAN - COMPLEMENT C1R	COMPONENT PRECURSOR				Novel Protein sim GBank	Ailagtatgtienipsessingt1 RAT - UDP-N-	ACETYLGLUCOSAMINE PEPTIDE N-	ACETYLGLUCOSAMINYLTRANSFERASE 110 KD	3	Novel Protein sim. GBank	gil4959442 gb AAD34351.1 AF12136 - (AF121360) DN/2DHHC/NEW1 zinc finger protein 11 [Drosophila	melanodaster]	13978464 (AF085693)	protein-coupled receptor kinase-associated ADP	ribosytation factor GTPase-activating protein [Rattus	norvegicus]				j							94232573 (3547, 3548) Novel Protein sim. GBank	gjj2495689JspjQ15034JY032_HUMAN - HYPOTHETICAL IPBOTEIN KJAA0032	
1766 95081201 (353 1763 80253216 (353 1769 87386888 (353 1770 95413144 (355 1771 94233542 (35 1772 87643510 (35 1773 94116824 (35	95081201 (3531, 3532) h	19200 (2001) 2006		 87755998 (3533, 3534)			87388988 (3537, 3538)		(2002, 2003)				·	_	יביים (יביים)				87643510 (3543, 3544)			_				-										1774 94232573 (3547, 3548)		

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65274572, 56182575, 56994075, 35696286, 22278997, 29331822, 29331828, 60432289, 29331828, 35696286, 22378997, 29331828, 2643289, 29331828, 5612502, 264828, 56182435, 264511, 265007, 265009, 60170031, 6043229, 60433438, 55812038, 21906754, 28968842, 87188559, 264601, 265017, 265018, 265019, 264762, 264448, 265022, 264691, 3957065, 21906765, 21906765, 21906767, 21906768, 265020, 265021, 265022, 264691, 3365703, 264632, 264634, 264638, 264634, 264638, 264634, 264638, 264634, 264638, 264488, 264488, 264632, 264634, 264638, 264488, 66326486, 87168518, 60432113, 22279000, 22279002, 264482, 264486	264910	56994075, 29331826, 265008, 87168474, 265017, 265018, 264761, 55811150, 264764, 56181562, 264689, 21906765, 21906768, 21906769, 33657023, 35695763, 60431528, 35696423, 55811576, 35695855, 22279000, 22279002, 284564	29331826, 29331827, 35696052, 264512, 265007, 265009, 265017, 265019, 264762, 18108351, 264769, 21906765, 21906766, 21906767, 21906767, 21906769, 265020, 265021, 264691, 264693, 18108370, 264556, 83373044	29148627, 35696286, 29147620, 265006, 265007, 265008, 18108385, 65274727, 264482, 264369, 264766	264107, 33657109, 56526486	264508, 264906, 264639	264259, 28331822, 264508, 264905, 264906, 264908, 264907, 264907, 265009, 264910, 264591, 264591, 264768, 264764, 264288, 264768, 264769, 264563, 264635, 264637, 264639, 264553	264768
UNCLASSIFIED		nucl_recpt	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED		interferon
			·		Contains protein domain (PF00807) - UNCLASSIFIED Apidaecin			
95359330 (3549, 3550) Novel Protein sim. GBank gil 1469199jdbj BA408487  - (050928) The KIAA0138 gene product is novel. [Homo sapiens]	94133756 (3551, 3552) Novel Protein sim. GBank gil4589676 dbj BAA76857.1  - (AB023230) KIAA1013 protein [Homo sapiens]		94851624 (3555, 3556) Novel Protein sim. GBank gil3875648[emb]CAA91454.1] - (Z66561) Similarity to Human rab13 protein (PIR Acc. No. A49647). Contains the ATP/GTP-binding site motif (PROSITE PS00017); cDNA EST EMBL:M89412 comes from this gene; cDNA EST yk212g9.3 comes from this gene; cDNA EST yk212g9	94133758 (3557, 3558) Novel Protein sim. GBank gil4589676 dbj BAA76857.1  - (AB023230) KIAA1013 protein [Homo sapiens]			88094607 (3563, 3564) Novel Protein sim. GBank gij729225 sp P41237 CTXN_RAT - CORTEXIN	1783   85717905 (3565, 3566) Novel Protein sim. GBank gi 2257543 dbj BAA21436  • (AB004538) protein arginine N-methyltransferase [Schizosaccharomyces pombe]
95359330 (3549, 3550)			8   94851624 (3555, 3556)			1 84047477 (3561, 3562)	2   88094607 (3563, 3564)	3  85717905 (3565, 3566)
1775	1776	1771	1778	1779	1780	1781	1782	178

35696286, 264259, 35696052, 264508, 264905, 264905, 264907, 68112502, 264908, 264905, 264907, 68112502, 264908, 264909, 265007, 265009, 265009, 265009, 264910, 265001, 265001, 264002, 264004, 265001, 264001, 264002, 264004, 264005, 1810831, 264763, 264768, 264631, 264691, 264691, 264691, 264691, 264691, 264691, 264692, 264693, 264632, 264633, 18108385, 2279000, 22279002, 264555, 264656, 264686	65274572, 264259, 29531022, 25531024, 29331825, 29331827, 29331828, 264905, 264906, 264908, 66712502, 56182435, 264511, 265007, 6043336, 55811150, 264683, 284369, 264687, 52644229, 21906767, 52644150, 33657023, 65274620, 33857182, 65274791, 35695855, 264555, 65274727, 22279002	264908, 35696423, 264636	264488, 264905, 264908, 264909, 264599, 264764, 264766, 264692, 60431528, 264629, 264636, 264564, 264566	264488, 83373044 264488, 29331828, 264909, 18108351, 264288, 265021, 264555, 264636	35696052, 264905, 264906, 264907, 264909, 264909, 265008, 264910, 264758, 265011, 265019, 264766, 264769, 264628, 264635
struct	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	
1755049 (U55042) - myosin X Contains protein domain (PF00169) - struct	·				
95197093 (3567, 3569) Novel Protein sim. GBank gil1755049 (U55042) - myosin X [Bos taurus]	95357475 (3569, 3570) Novet Protein sim. GBank gil4589552 dbj BAA76798.1 - (AB023171) KIAA0954 protein (Homo sapiens)	85296465 (3571, 3572) Novel Protein sim. GBank gi[117788 sp P26770 CYA4_RAT ADENYLATE CYCLASE, TYPE IV (ATP PYROPHOSPHATE-LYASE) (ADENYLY CYCLASE)	87434784 (3573, 3574) Novel Protein sim. GBank gij3877175 emb CAA90338.1 - (Z50028) cDNA EST yk321h8.5 comes from this gene; cDNA EST EMBL:D68896 comes from this gene; cDNA EST yk395f9.5 comes from this gene (Caenorhabdilis		
15197093 (3567, 3568)	95357475 (3569, 3570)	85296465 (3571, 3572)	87434784 (3573, 3574)		82489734 (3579, 3580)
1784	1785	1786	1787	1788 1789	1780

264488, 264686, 264687, 264768, 18108394, 264769, 18108397, 264259, 264691, 264692, 33557023, 264693, 264593, 264692, 264628, 264907, 264629, 264908, 264909, 264510, 265009, 264901, 264632, 264630, 264631, 264631, 264631, 264632, 264637, 264635, 264536, 264636, 264637, 264637, 264638, 18108381, 264639, 264567, 264604, 264760, 264604, 264761, 264681, 264762, 264583, 264763, 264681	13657402, 87168559, 264369, 33657023, 3569855, 20281071, 264559, 18108387, 87168559, 18108387, 8716858	052/4974, 227,0393, 227,0394, 227,0399, 264093, 264259, 29331824, 264103, 264105, 29331830, 264103, 264105, 29331830, 266007, 264910, 265009, 60170831, 60433356, 21906754, 265010, 265017, 265019, 264681, 264682, 21906767, 21906768, 21906769, 265020, 265021, 265022, 60170615, 52644150, 33657109, 18108370, 18108374, 65274791, 20281071, 60432113, 22278000, 264482, 264564	18108394, 222,18399, 222,16399, 204239, 29331822, 29331824, 29331825, 29146498, 29316492, 264508, 264905, 52644045, 264112, 265006, 265008, 264910, 60433356, 264712, 265017, 1910351, 24763, 26448, 264683, 265369, 21906765, 21906766, 21906769, 21906769, 21906769, 21906769, 21906769, 21906769, 21906769, 2164538, 264529, 18108374, 18108378, 35696423, 35695855, 264556, 264557, 264638, 264558, 18108374, 18108378, 264638, 264558, 18108385, 264554	264632, 264635, 264636, 264595, 264396, 264907, 264566, 264909	264488, 264907, 264909, 264594, 264595, 264766, 264687, 21906765, 21906767, 264628, 264630, 264559
	UNCLASSIFIED	A i Pase_associated	UNCLASSIFIED	UNCLASSIFIED	glycoprotein
Contains protein domain (PF00047) - giycoprotein Immunoglobutin domain	Contains protein domain (PF01585) - UNCLASSIFIED G-patch domain		Contains protein domain (PF01798) - UNCLASSIFIED Putative snoRNA binding domain		
. 5	87792690 (3583, 3584) Novel Protein sim. GBank gil4337106 gb AAD18082 - (AF129756) BAT4 [Homo sapiens]	95337877 (3585, 3586) Novel Protein sim. GBank gil5579331[gb]AAD45504.1 AF14573 - (AF145732) endoplasmic reticulum alpha-mannosidase 1 [Homo sapiens]	88) Novel Protein sim. GBank gil4914604 emb CAB43677.1 - (AL050369) hypothetical protein [Homo sapiens]	(085)	86599486 (3591, 3592) Novel Protein sim. GBank gij585084 sp Q07803 EFGM_RAT - ELONGATION FACTOR G, MITOCHONDRIAL PRECURSOR (MEF-G)
1781 95197259 (3581, 3582 <u>)</u>	1792 87792690 (3583, 3584	1783 95337877 (3585, 3586)	1794 87759806 (3587, 3588)	1795 79747856 (3589, 3590)	1796 86599486 (3591, 359

22278996, 22278997, 22278998, 22278999, 29331822, 264910, 60170831, 21906754, 52644229, 21906768, 21906768, 21906769, 35695917, 265022, 52644150, 264691, 33657023, 263967, 33657109, 22279000	22278994, 56994075, 22278997, 22278998, 22278999, 264259, 29331826, 6043229, 29331826, 33556970, 265008, 6043229, 264757, 60433438, 21908754, 33657084, 87168559, 285017, 18108351, 284682, 285017, 18108351, 284682, 21908768, 21908768, 21908769, 35695917, 265020, 265021, 33657023, 33657182, 27486261, 27486265, 33657349, 263973, 18108374, 55811576, 35695855, 18108385, 87168518, 22279000, 264486	264908, 21906754, 21906767, 21906769, 265020, 33657023, 264692, 264693, 264404, 22279000	264691, 264556, 264566	52644045, 265007, 264632	35696286, 66714117, 264508, 264509, 56182435, 264512, 18108351, 264686, 55811957, 264692, 55811576, 35695855, 284486	264766, 264488, 264687, 264489, 264766, 264769, 264769, 264689, 21906769, 35696286, 35695917, 264691, 264691, 264692, 246693, 20281099, 18108364, 35696052, 264508, 264509, 264509, 264905, 18108374, 18108376, 35696423, 35695855, 264510, 264511, 265006, 265007, 264512, 265008, 264910, 264631, 264637, 264592, 264634, 264592, 264636, 264591, 264636, 264037, 264592, 264636, 264634, 264592, 264592, 264594, 264592, 264594, 264592, 264594, 264592, 264594, 264592, 264594, 264592, 264594, 264592, 264594, 264592, 264594, 264592, 264502, 264592, 264592, 264592, 264592, 264592, 264592, 264592, 264502, 264592, 264592, 264592, 264592, 264592, 264592, 264592, 264502, 26
ribosomalprot	peptidase	qdə	UNCLASSIFIED		·	
GBank gi 1842111 (U87586) - decoy	Novel Protein sim. GBank gi 2832906 dbj gAA24608.1  - (D89340) dipeptidyl peptidase III [Rattus norvegicus]	GBank gi 5689541 db  BAA83054.1  - 102 protein [Homo sapiens]		GBank 27729.1 AF13295 - (AF132954) CGI-20 ens}	GBank 37 SSGP_VOLCA - SULFATED PROTEIN 185 (SSG 185)	GBank 27729.1 AF13295 - (AF132954) CGI-20 ens}
Novel Protein sim. GE [Arabidopsis thaliana]	Novel Protein sim. ( (D89340) dipeptidyl	Novel Protein sim. GBank g (AB029025) KIAA1102 prot	-	Novel Protein sim. GBank gil4680679 gb AAD27729. protein [Homo sapiens]	Novel Protein sim. GBank gi 134920 sp P21997 SSGP SURFACE GLYCOPROTEI	Novel Protein sim. GBank gi 4680679 gb AAD27729. protein [Homo sapiens]
1797   91223219 (3593, 3594) Novel Protein sim. GBank [Arabidopsis thaliana]	91221276 (3595, 3596) Novel Protein stm. GBank g (D89340) dipeptidyl peptids		87080116 (3599, 3600)	95060723 (3601, 3602)	87771012 (3603, 3604)	95060725 (3605, 3606) Novel Protein sim. GBank , gil46806791gbJAAD27729, protein [Homo sapiens]
1797			1800			1803

					300736 30001000 10001000 1000
88	97770203 (3607, 3608)	1804 87770203 (3607, 3608) Novel Protein sim. GBank gil3879914 emb CAA98538.1 -			52646365, 22276997, 22276999, 204905, 564908, 564909, 564910, 21906754, 264766,
		(Z74043) predicted using Genelinder; cUNA ES1			21906764 21906768 35695917 265020
_		EMBL:C13850 comes from this gene; cDNA EST			21500103, 21500100, 33035311, 200213,
	<del></del> -	EMBL:C11575 comes from this gene; cDNA EST yk343f4.5			203022, 204031, 204037, 204033, 22213000, 1
		comes from this gene (Caenorhabditis elegans)			204304, 204300
805	95330375 (3609, 3610)	95330375 (3609, 3610) Novel Protein sim. GBank			29331824, 29331825, 29331826, 29331827,
		gij5453644 ref NP_006461.1 pEBBP - estrogen-responsive	•		29331828, 87,108339, 204286, 204887,
					52644229, 33696423, 264636, 60432113
1806	94133762 (3611, 3612)	94133762 (3611, 3612) Novel Protein sim. GBank gil4589676 dbj BAA76857.1  -	18	struct	264094, 264105, 264908, 33696423, 263006.
		(AB023230) KIAA1013 protein [Homo sapiens]			265007, 265008, 264555, 264582, 265011,
					20224024 284000 284040 43647023
1807	86943032 (3613, 3614)				2933 1024, 204300, 204310, 33037 023,
	10000 4000 00000	11 3000 March 20140040701 Jan 20 20 20 20 20 20 20 20 20 20 20 20 20		UNCLASSIFIED	284488, 35696286, 66714117, 35696052,
1808	(3195, 3613) 117249/8	87642711 (3615, 3616) Novel Protein Sim. GBank gildso4079 einb CA0432311  *	•		66712502, 264592, 60433438, 52644296.
		(ערספססס וואלאסוויפויסן לאיסיים וואלאסוויפויסן איסיים איסי			265010, 264683, 264369, 284689, 55811957.
					35695917, 33657109, 35695763, 55810764,
					18108379, 35696423, 35695855, 56182323.
					264563, 264564, 264487
	10100 11000 001 001	S ordinates (190050) // (190050) - 190050 - 190050   190050		UNCI ASSIFIED	264594, 55811150, 264686, 29148629,
5001	83321400 (3017, 3010)	<u> </u>			29148784, 264690, 264629, 18108374,
		protein (mus muscalus)			264556, 264557, 264558
1010	88006218 /2810 3820N	88006318 (3810 3820) Navel Protein eim CBank		UNCLASSIFIED	264488, 35696052, 264905, 264906, 264907,
_	0202 (2018, 2020)	nitras2944tenip471791Y.19P YEAST - HYPOTHETICAL			264908, 264909, 264511, 265009, 264910,
		118 4 KD DEOTEIN IN BAT2. DAI 5 INTERGENIC REGION			264592, 264593, 264594, 33657402, 264757.
	j				264595, 264758, 264596, 264759, 264600.
		10000000000000000000000000000000000000			264601, 264762, 264683, 264764, 264288,
	,				264684, 264766, 264767, 264686, 264768.
	•				264687, 264769, 264689, 265021, 264690.
					264691, 264693, 264628, 264629, 18108374.
					264630, 264631, 264632, 264634, 264635,
					264636, 264637, 264638, 264639, 264563.
					264566, 264486, 264567
1811	IRRORG272 (3821 3622)	88086272 (3621 3622) Novel Protein sim GBank gil2134984birll37275 - death-	Contains protein domain (PF00023) - kinase	kinase	264488, 264259, 264508, 264509, 264905.
	(======================================	(EC 2.7.1-) - human	Ank repeat		264906, 264907, 56182435, 264511, 264512.
					264910, 264758, 265011, 264600, 264604,
					18108354, 264766, 264686, 264769, 264534,
					60170615, 33657023, 264629, 264631,
					264639, 264563, 264482, 264483
1812	79245772 (3623, 3624)				29331822, 29331824, 265019, 18108351,
					21906769

772 (3625, 3626) 747 (3627, 3628) 173 (3629, 3630) 345 (3631, 3632)	1813 88090972 (3625, 3628) Novel Protein sim. GBank gil3843608 (AC005395) EH domain-binding mitotic phosphoprotein [Homo sapiens] domain-binding mitotic phosphoprotein [Homo sapiens] hypothetical protein sim. GBank gil3843608 (AC005395) - hypothetical protein [Arabidopsis thaliana] hypothetical protein sim. GBank gil117788 sp P26770 CYA4_RAT ADENYLATE CYCLASE, TYPE IV (ATP PYROPHOSPHATE-LYASE) (ADENYLYL CYCLASE) (ADENYL CYCLASE) (ADENYLYL CYCLASE) (ADENYLYL CYCLASE) (ADENYLYL CYCLASE) (ADENYLYL CYCLASE) (ADENYLYL CYCLASE) (ADENYL CYCLASE)	Contains protein domain (PF01417) - glucoamylase ENTH domain UNCLASSIFIE	g g	56182575, 284259, 29331824, 66714117. 29331828, 35696052, 284509, 264905, 264907, 264907, 264908, 66712502, 264909. 265007, 264910, 264591, 264593, 55812038, 265007, 264910, 264591, 264593, 264764, 264683, 264769, 264682, 264764, 264683, 264769, 264681, 264769, 264683, 264769, 264634, 264635, 26279002, 264639, 264634, 264635, 26279002, 264636, 264634, 264635, 26279002, 264268, 26279002, 264268, 26279002, 264269, 29331824, 29331824, 29331825, 69331824, 26331826, 29331826, 29331826, 29331826, 29331826, 29331826, 264968, 26409, 264010, 265011, 265021, 265021, 265022, 2643999, 264599, 2645907, 264908, 264909, 265001, 264766, 264691, 265011, 264760, 264591, 265011, 264591, 265022, 264691, 18108394, 18108394, 18108397, 264599, 264591, 265011, 264564, 264564, 264567, 264567, 264564, 264567, 264567, 264564, 264567, 264567, 264564, 264567, 264564, 264567, 264567, 264564, 264567, 264567, 264564, 264567, 264567, 264564, 264567, 264567, 264564, 264567, 264564, 264567, 264
3634)	8809526B (3633, 3634) Novel Protein sim. GBank gil3766377 emb CAA21429  - (AL031907) hypothetical protein [Schizosaccharomyces pombe]	Contains protein domain (PF00400) - histone WD domain, G-beta repeat		264488, 22278997, 22278999, 60432049, 29331822, 29331824, 60432289, 52644045, 60170831, 265017, 265018, 265019, 18108351, 264682, 52644229, 21906765, 21906767, 21906768, 52644150, 33657023, 33657109, 27486262, 18108370, 18108374, 60170394, 56182323, 22279002
1636)	85806775 (3635, 3636) Novel Protein sim. GBank gij3879121[emb CAA94370] - (Z70310) predicted using Genefinder; Similarity to Mouse ankyrin (PIR Acc. No. S37771); cDNA EST EMBL:T01923 comes from this gene; cDNA EST EMBL:D32335 comes from this gene; cDNA EST EMBL:D32723 comes from this gene; cDNA EST EMBL:D32723 comes from this	Contains protein domain (PF00023) - transcriptfactor Ank repeat		35696286, 60433356, 26458, 264369, 264686, 21906769, 264693, 264632
1638)	87759572 (3637, 3638) Novel Protein sim. GBank gil5031865 ref NP_005771.1 pLHFP - lipoma HMGIC fusion partner		UNCLASSIFIED	22278995, 22278997, 22278999, 264259, 29331826, 264908, 265007, 265008, 265009, 60432229, 265017, 265018, 265019, 26448, 264288, 21906768, 21906769, 265020, 18108381, 18108384, 22279000, 22279002, 264567

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264905, 264907, 264594	264907, 264768, 263978	22278995, 56994075, 22278996, 22278997,	22278998, 264259, 29331824, 29331825,	29331826, 35696052, 29331828, 264908,	29331830, 60170831, 264591, 264593,	60433356 264598 265017, 265019.	18108351 264763 264683 21906765	21908281, E011081, E0	265020, 265021, 33657023, 18108364,	18108370, 35695855, 22279000, 22279002	264488, 264259, 264511, 264288, 264768,	264693, 35696423, 264634, 18108385,	264486	264907, 264908, 264909, 264511, 264631.	264634, 264635, 264637, 264638, 264639,	264758, 264566	60432049, 264910, 264487	COLCOO CICCION COLON	ATPase_associated [52644507, 52645156, 52646844, 22276994,	22278996, 56994075, 264259, 60432049,	52645080, 35696052, 66712502, 52644045,	265008, 265009, 60432229, 60433356,	60433438, 52646317, 52644298, 265011	87168559, 264448, 264288, 264369, 264688.	52644229, 264689, 21906765, 21906768,	265020, 60170615, 52644150, 33657023,	27486262, 27486264, 27486265, 35695763,	35696423, 35695855, 83373044, 87168518,	264404, 22279002	
											UNCLASSIFIED						UNCLASSIFIED		ATPase_associated											
														Contains protein domain (PF00047) - UNCLASSIFIED	Immunoqlobulin domain	,														
		01221573 (3643 3644) Novel Protein sim GBank nil4884130lemblCAB43272.11												BER12025 (3547 3548) Novel Protein sim GBank pil477072lpirilA48018 - mucin 7			87430125 (3649, 3650) Novel Protein sim. GBank gij3036803 emb CAA18493  -	(AL022373) hypothetical protein [Arabidopsis thallana]	91723612 (3651, 3652) Novel Protein sim. GBank	ail4880885labIAAD27732.11AF13295 - (AF132957) CGI-23	protein (Homo saniens)									
87769455 (3839 3840)	80431510 (3841 3642)	01221523 (3643 3644)	1,221,221,221,531,5								OCESSON (SEAR SEAR)	0222230 (2043, 3040)		86612025 (3647 3648)	מספו להספי (מספו מספו		_	,	91723612 (3651, 3652)										*	
1820	_		770									1953		183	1064		1825		1826				_		_					

65274572, 29331822, 29331824, 29331825, 66714117, 29331826, 29331827, 264907, 264909, 5264909, 5264909, 5264909, 2654009, 265400, 265009, 265009, 265009, 265009, 265009, 265009, 265009, 265009, 264509, 33657023, 264693, 18108364, 33657109, 18108384, 264628, 55810764, 56182323, 18108384, 264564	264488, 56182435, 264769, 29331826, 29331828, 264511, 265006, 265007, 264910, 264631, 264509, 264690, 264636, 264564, 264691, 60432229, 60432049, 264259, 264629, 33657023, 264486, 264909, 264567, 264595, 284766	264908, 265022, 33657023, 87168518, 22279002	264259, 29331824, 264907, 264908, 66712502, 264510, 265007, 265008, 55812038, 265018, 21908765, 52644150, 33657109, 264555, 264556, 264557, 56182323, 18108382, 83373044, 18108385, 264564	264905, 264908	22278997, 264259, 29331824, 264909, 18108351, 263974, 22279002	29331825, 29331826, 29331827, 265017, 264683, 264288, 264766, 264768, 21906767, 21906768, 264692, 22279002	56182575, 265018 60432049, 264908	29331824, 35696052, 264910, 60433438, 264688, 35695917, 265020, 52644150, 65274620, 52644332
				UNCLASSIFIED	transcriptfactor	tm7	ATPase_associated	dna_ma_bind
1	·		Contains protein domain (PF00628) - transcriptfactor PHD-finger	Contains protein domain (PF00008) - UNCLASSIFIED EGF-like domain	Contains protein domain (PF00010) - transcriptfactor Helix-toop-helix DNA-binding domain		Contains protein domain (PF00008) - ATPase_associated EGF-like domain Contains protein domain (PF00169) - struct	Zinc finger, C2H2 type
90992645 (3683, 3684) Novel Protein sim. GBank gil 1326268 (US8728) - C54H2.1 gene product [Caenorhabdilis elegans]	0	87444764 (3687, 3688) Novel Protein sim. GBank gi[2496887]sp Q09232 YQ22_CAEEL - HYPOTHETICAL 32.0 KD PROTEIN C09F5.2 IN CHROMOSOME III	95096673 (3689, 3690) Novel Protein sim. GBank gij 1175494 sp Q09819 YAC5_SCHPO - HYPOTHETICAL 45.0 KD PROTEIN C16C9.05 IN CHROMOSOME I	84287872 (3691, 3692) Novel Protein sim. GBank gij3881080jembjCAA21739j - (AL032657) similar to EGF-fike domain; cDNA EST yk299a12.3 comes from this gene; cDNA EST EMBL:D35398 comes from this gene; cDNA EST yk331h8.5 comes from this gene; cDNA EST yk299a12.5 comes from this gene; cDNA EST yk299a12.5 comes from this gene; cDNA EST yk299a12.5 comes from	87821487 (3693, 3694) Novel Protein sim. GBank gij5059323jgbJAAD38967.1JAF15162 - (AF151522) hairy and enhancer of split related-1 (Homo sapiens)	1848 86789360 (3695, 3696) Novel Protein sim. GBank gil5701854 emb CAB52191.1  - (AJ245417) G5b protein [Homo sapiens]	84287974 (3697, 3698) Novel Protein sim. GBank gi[4503665[ref]NP_001989.1]pFBLN - fibulin 2 precursor 86689650 (3699, 3700) Novel Protein sim. GBank gi[4589562[db]]BAA76813.1] -	[AB023189] (3701, 3702) Novel Protein sim. GBank gi[220637]dbj[BAA01477] - (D10627) zinc finger protein [Mus musculus]
	95292692 (3685, 3686)	87444764 (3687, 3688	95096673 (3689, 3690			86789360 (3695, 3696		
1842	1843	1844	1845	1846	1847	<u>\$</u>	1849 1850	1851

	UNCLASSIFIED 224687, 264768, 52644507, 264769, 21906768, 21906767, 21906768, 22278995, 56994075, 22278999, 5264150, 264259, 26994075, 22278999, 5264150, 264259, 264692, 29331822, 29331824, 52645129, 29331827, 3365970, 33657349, 35695763, 264690, 266906, 264900, 266904, 264512, 264630, 265009, 265006, 264514, 264639, 264630, 265009, 264631, 264631, 264634, 264639, 265437, 264539, 264634, 264639, 265437, 264539, 264632, 264639, 264438, 264662, 264682, 264438, 265018, 264766, 264567, 264288, 264369, 264369, 264766, 264567, 264288, 264369, 264766			UNCLASSIFIED 56994075, 264259, 29331828, 264511, 264910, 264758, 264693, 264637, 18108381, 83373044	ATPasa_associated   22278999, 264259, 264906, 60170831,   264448, 264686, 265020, 265022, 33657109,
UNGLA	UNCLA	nuclease	Contains protein domain (PF01344) -  nucl_recpt Kelch motif	חאכת	ATPas
Novel Protein sim. GBank gi[5174629fæf]NP_006090.1[pPIAS - protein inhibitor of activated STAT3		gi 2072964 (U93569) - putative	gil4539520 emb CAB39994.1  - ovel protein similar to Drosophila KEL) and a heterogenous set of omo sapiens]	94231871 (3711, 3712) Novel Protein sim. GBank gi[3954978 emb CAA06945  - (AJ008278) acetylglucosaminyltransferase-like protein [Mus musculus]	94324455 (3713, 3714) Novel Protein sim. GBank gil4322670[gb]AAD16120] - (AF094508) dentin phosphoryn (Homo saplens)
		1854 86038152 (3707, 3708)		1856  94231871 (3711, 3712)	1857   84324455 (3713, 3714)

264757	22278996, 29331824, 265007, 33109954, 265019, 264369, 21906768, 29148784, 27486261, 52644332, 22279002	265019	264887, 264259, 28331822, 28331824, 29331825, 265007, 265009, 264581, 33109954, 265010, 265019, 264369, 264288, 264686, 264691, 264693, 27486284, 18108370, 18108374, 263977, 55811576, 26182323, 264639, 22279000, 22279002, 264482	35696286, 264259, 87168474, 264369, 21906766, 264558, 264563	264601, 264766, 29148627, 29148629, 264692, 264629, 264635	22278999, 264259, 264807, 265018, 18108370, 264634, 264635, 264555, 264556. 264638, 18108387	22278999, 264490, 29331822, 6674317, 65712502, 265006, 265007, 265008, 265009, 264591, 60433438, 265010, 265019, 264760, 264448, 264768, 29146627, 29148629, 265020, 265022, 18109385, 60432113	18108374, 264789, 18108377, 21906765, 21906768, 35696423, 56182575, 21908769, 21906768, 35696423, 56182575, 21908769, 29148629, 3569586, 35695917, 2545021, 264510, 264512, 264512, 264552, 264553, 264556, 264659, 264557, 33657023, 6043336, 29331822, 264559, 264559, 264559, 264559, 264559, 264559, 264559, 264559, 264557, 33657182, 29331827, 35696052, 33656970, 87168518, 265017, 60431602, 22279000, 264508, 264509, 18108351, 264966	264094
				kinase	UNCLASSIFIED	synthase	·	rinase	UNCLASSIFIED
Contains protein domain (PF00312) - inbosomalprot Ribosomal protein S15			Contains protein domain (PF00285) - Citrate synthase	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain		Contains protein domain (PF00754) - synthase F5/8 type C domain	·		
87628311 (3715, 3716) Novel Protein sim. GBank gil4981903jgbJAAD36415.1JAE00178 - (AE001788) gilvecomel protein S15 (Themploos marifima)	84407464 (3717, 3718) Novel Protein sim. GBank gil4240317[dbj BAA74937.1] - (AB020721) KIAA0914 protein [Homo saplens]		88086370 (3721, 3722) Novel Protein sim. GBank gil2143637[pir]  845,05 - calcium-dependent actin-binding protein - rat	87372923 (3723, 3724) Novel Protein sim. GBank gij125493 sp P07313 KMLC_RABIT - MYOSIN LIGHT CHAIN KINASE SKELETAL MUSCLE (MLCK)	85775037 (3725, 3726) Novel Protein sim. GBank gij3820909 emb CAA09299  -	85547832 (3727, 3728) Novel Protein sim. GBank gil4322263 gblAAD15985  - (AF077738) metallocarboxypeptidase CPX-1 [Mus muscutus]	1865 87740827 (3729, 3730) Novel Protein sim. GBank gilz495727 sp Q93073 Y256_HUMAN - HYPOTHETICAL PROTEIN KIAA0256	87266816 (3731, 3732) Novel Protein sim. GBank gil5262617 emb CAB45748.1  - (AL080157) hypothetical protein [Homo saplens]	84579159 (3733, 3734) Novel Protein sim. GBank gij3859930 (AF078096) - forkhead/winged helix-like transcription factor 7 [Homo sapiens]
1858 87628311 (3715, 3716	1859 84407464 (3717, 3710	1860 17929308 (3719, 372	1861   88066370 (3721, 372	1862 87372923 (3723, 372	1863 85775037 (3725, 372	1884 85547832 (3727, 372	865 87740827 (3729, 37.	1866 87266816 (3731, 37	1867 84579159 (3733, 37

	264489, 22278997, 22278999, 29331825, 29331826, 265008, 265009, 33657402, 87168474, 18108351, 21906765, 21906768, 21906769, 265020, 265021, 60170615, 27486264, 264628, 18108374, 264631, 18108385, 87168518, 22279000, 22279002, 264568, 264567	264508, 264509, 264906, 264908, 264909, 264910, 264591, 264600, 18108351, 264683, 264766, 264769, 35695855, 264634, 264558, 264639, 18108385, 264563, 264486	29331824, 60424269, 265007, 265008, 21906754, 265017, 265018, 265019, 264288, 264766, 264686, 264688, 21908768, 21906769, 35695917, 60170615, 264692, 18108368, 35695763, 35696423, 65274791, 264638, 264639, 56526486	264509, 264905, 264595, 264768, 264635, 264636, 264563, 264486	264510, 264594, 264565	264509, 264512, 265009, 265011, 18108351, 264687, 264691, 18108370, 18108374, 264635	264488, 35695917, 264259, 264905, 264907, 264908, 264909, 263978, 264511, 264635, 264636, 264638, 33857402, 264558, 18108385, 264600, 264604, 264764, 264567, 264766	35696286, 264828, 264592, 264557, 264558	264259, 29331822, 60432289, 264908, 264909, 264604, 264784, 264288, 264769, 18108376, 264556, 264558, 264559,	264767, 264686, 264768, 263978, 264693, 264639, 264639, 265010, 264563, 264905, 264906, 264907
gij3881525[emb]CAA938841:D32579 comes from this gene; comes from this gene; condes from this gene; condes from this gene; condes from this gene; cDNA in this gene; cDNA EST is gene [Caenorhabditis elegans] sene [Caenorhabditis elegans] is gene [Caenorhabditis elegans] ent protein [human herpesvirus 2] ent protein [human herpesvirus 2] ent protein [human herpesvirus 2] is gij2460316 (AF022147) - uterus-tsmembrane protein [Rattus is gij4589520[dbjjgAA76782.1] - tein [Homo saplens] gij263810[bbs[122920 - collagen illa=tube worms, Peptide, 1027 ilgj3983356[gbjAAC83924.1] - ing protein precursor [Scaphiopus ing protein precursor [Scaphiopus]	nuclease	glycoprotein	UNCLASSIFIED	UNCLASSIFIED	profease		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED
		Contains protein domain (PF00560) - Leucine Rich Repeat				•	Contains protein domain (PF00293) - Bacterlal mutT. protein			
	1868 87357459 (3735, 3735) Novel Protein sim. GBank gil3881525 emblCAA93884 - (Z70038) cDNA EST EMBL:D32579 comes from this gene; cDNA EST EMBL:D35254 comes from this gene; cDNA EST yk224b3.5 comes from this gene; cDNA EST yk357f10.5 comes from this gene [Caenorhabditis elegans]				Novel Protein sim. GBank ovary specific putative tran norvegicus]	3745, 3746)	3747, 3748)	3749, 3750) Novel Protein sim. GBank gil4589520 dbj BAA76782.1  - (AB023155) KIAA0938 protein [Homo sapiens]	3751, 3752) Novel Protein sim. GBank gil263810jbbs 122920 - collagen alpha chain [Rifiia pachyptila=tube worms, Peptide, 1027 aa]	

264488, 18108394, 56181686, 22278998, 60432049, 264259, 29331822, 29331824, 29331825, 29331824, 29331825, 29331827, 35696052, 264508, 264509, 264905, 264905, 264907, 264509, 264909, 264907, 264909, 264909, 264907, 264501, 264510, 264511, 264503, 264510, 264510, 264513, 264500, 265011, 87168559, 264600, 265017, 264604, 265019, 264605, 264680, 264761, 294862, 264769, 264680, 264681, 33657023, 264682, 264693, 33657109, 18108370, 264628, 60431528, 264634, 60431850, 264637, 264634, 60431850, 264637, 264634, 264632, 264631, 264634, 264639, 264631, 264634, 264639, 264631, 264632, 263373044, 181083385, 264539, 264537, 264633, 264634, 264565, 264566, 264537, 264634, 264564, 264565, 264566, 264537, 264634, 264564, 264565, 264566, 264537, 264634, 264564, 264565, 264566, 264537, 264634, 264564, 264565, 264566, 264537, 264564, 264564, 264565, 264566, 264565, 264567, 264567, 264567, 264567, 264507, 2645	264596, 264766, 265020, 264628, 80431528, 264634, 56526486, 264080, 284563			263978	264906, 55612038, 264756, 255010, 255019, 265019, 18108351, 264286, 21906765, 21906768, 21906769, 265021, 33657023, 33657109, 56182323, 83373044, 18108385, 22279000, 22279002	18108398, 29147620, 264907, 265009, 264600, 265018, 18108351, 264288, 264689, 21908765, 21906768, 21908769, 264691, 264692, 264693, 264628, 18108370, 264536, 264558, 264404
	UNCLASSIFIED		ATPase_associated		heilcase	UNCLASSIFIED
5			Contains protein domain (PF00168) - ATPase_associated	Contains protein domain (PF01454) - MAGE family	Contains protein domain (PF00176) - helicase SNF2 and others N-terminal domain	
1884   95310885 (3767, 3768)  Novel Protein sim. GBank gj[4929643]gbpAD34082.1 AF15184 - (AF151845) CGI-87 protein [Homo sapiens]	87644280 (3769, 3770) Novel Protein sim. GBank gi 2507155 sp P37370 VRP1_YEAST - VERPROLIN	86674062 (3771, 3772) Novel Protein sim. GBank gij2854158jgbjAAC02577.1  - (AF045641) No definition line found [Caenorhabditis elegans]	olein sim. GBank 21 ref NP_006023.1 pCPNE - copine VI (neuronal)	1888 87822804 (3775, 3776) Novel Protein sim. GBank gij3319931jembjCAB108411 - (Z98046) dJ14O9.2 (Melanoma-Associated Antigen MAGE	91255783 (3777, 3778) Novel Protein sim. GBank gil1083308 pir  A56559 - enhancer-trap-locus-1 protein - mouse (fragment)	87626705 (3779, 3780) Novel Protein sim. GBank gil4240195 dbj BAA74876.1  - (AB020660) KIAA0853 protein [Homo sapiens]
95310885 (3767, 376)	1885 8764280 (3769, 377	1886   86674062 (3771, 377	1887 94139139 (3773, 37	888 87822804 (3775, 37	1889   91255783 (3777, 37	1890   87626705 (3779, 37

	22278995, 264509, 87168559, 18108351, 264448, 264682, 265020, 264683, 18108374, 22279000	264569, 65274572, 22278997, 22278999, 264259, 29331822, 29331824, 66714117, 29331826, 264906, 265006, 265008, 265009, 264592, 265018, 264681, 264448, 264683, 18108354, 264369, 21906768, 264685, 264766, 264687, 264689, 21906768, 265020, 265022, 60170515, 52644150, 264690, 264691, 284692, 33657029, 264693, 33657109, 264632, 264634, 264557, 264530, 264632, 264634, 264557, 264530, 264632, 264634, 264557, 264539, 264631, 18108384, 264557, 264558, 60170394, 18108381, 18108385, 22279000	264508, 264908, 264908, 264907, 264594, 264684, 264690, 264692, 264630, 264635, 264538, 264639, 264563	56182575, 264259, 60432289, 29331826, 264107, 264908, 264908, 264910, 60170831, 264758, 265010, 265018, 264448, 284288, 264768, 33657109, 264628, 55810784, 18108379, 264634, 56182323, 56526486		264259	22278996, 22278998, 22278999, 29331824, 29331825, 60432289, 29331827, 35696052, 29331828, 265008, 265019, 264681, 264682, 264448, 264369, 52644229, 21906765, 21906768, 21906769, 60170615, 55810764, 22279000	264259, 264508, 264909, 60432229, 264769, 21908765, 21906769	
UNCLASSIFIED	struct	laminin			UNCLASSIFII	cadherin	helicase	glycoprotein UNCLASSIFIED	
	Contains protein domain (PF01412) - is Pulative GTP-ase activating protein for Arf			Contains protein domain (PF00435) - Spectrin repeat	Contains protein domain (PF00627) - UNCLASSIFIED UBA domain			Contains protein domain (PF00909) - glycoprotein Ammonium Transporter Family UNCLASSIF	
	87642825 (3783, 3784) Novel Protein sim. GBank gil5689535 dbj BAA83051.1  - (AB029022) KIAA1099 protein [Homo sapiens]			87631881 (3789, 3790) Novel Protein sim. GBank gil5262574 emb CAB45729.1  - (AL080133) hypothetical protein [Homo sapiens]		80565569 (3793, 3794) Novel Protein sim. GBank gif728836 sp P39193 ALU6_HUMAN - IIII ALU SUBFAMILY SP WARNING ENTRY IIII	87617637 (3795, 3796) Novel Protein sim. GBank gi 127560 sp P23249 MV10_MOUSE - PROTEIN MOV-10	86873097 (3797, 3798) Novel Protein sim. GBank gil2909819 (AF031548) - erythrocyte membrane glycoprotein Rh50 (Homo sapiens) 87641858 (3799, 3800) Novel Protein sim. GBank gil4102881 (AF017250) -	vitellogenin precursor (Oreochromis aureus)
1891 87013895 (3781, 3782)		88533826 (3785, 3786)			85673555 (3791, 3792)	80565569 (3793, 3794)			
1691	1892	<del> </del>		1895	1896	1897	1898	1900	

5	95196647 (3801, 3	3802)	1901 95196647 (3801, 3802) Novel Protein sim. GBank gij585959 sp P38378 S61A_RAT   CPROTEIN SEC61 ALPHA CONTINUE STIDINIT	Contains protein domain (PF00242) - Iransport DNA polymerase (viral) N-terminal		264488, 52644507, 52645156, 18108396, 52846365, 52846842, 18108397, 58182575, 22278894, 22278895, 56994075, 35696286,
				.,.		262259. 2227899. 264490. 60432049. 264259. 29331824. 29331825, 6845080. 29331824. 29331825, 69432289. 29331827, 29331828. 35696052. 33656970. 2946498. 264906. 264907. 29331830.
						264908, 52644045, 264909, 264112, 265006, 264512, 265008, 264910, 265009, 60170831, 26142229, 6043438
						55812038, 264758, 33109954, 21908754, 33557084, 252644296, 87166474, 265010, 2566713, 2756718, 2756719
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						21906769, 55811957, 35895917, 265020. 265021, 52644150, 18108362, 33657023,
					-	264693, 263967, 3365/109, 3365/162, 27486264, 33657349, 35695763, 18108370,
						18108376, 55811576, 35696423, 35695855.
						60170394, 83373044, 18108385, 87168518,
1902		3804)	80202013 (3803, 3804) Novel Protein sim. GBank gij4426613[gbjAAD20451] -		dna_ma_bind	60432113, 264564 264107, 263976
			(AF098798) SLM-1 [Mus musculus]			001100 00011000 00011000 001000
1903		. 3806)	Novel Protein sim. GBank gij3747107 (AF095741) - unknown [Rattus norvegicus]		UNCLASSIFIED	264259, 29331625, 29331627, 264508, 264907, 265008, 60170831, 60433356,
	-					60433438, 284759, 21906754, 264448, 264288, 265021, 265022, 33657023, 264693,
						55811576, 264555, 264556, 22279000
1904		, 3808)	80434213 (3807, 3808) Novel Protein sim. GBank		struct	264509, 264905, 264906, 264907, 264908,
			gil1352911jspjP47147jYJ80_YEAST - HYPOTHETICAL			265007, 264910, 264686, 264768, 264687. 264769, 264693, 264628, 18108374, 284634,
						264636, 264637, 264565
1905		, 3810)	95351140 (3809, 3810) Novel Protein sim. GBank gij3043714(dbjjBAA25521)	Contains protein domain (PF00293) -		264488, 264768, 264769, 264689, 29148629,     34604017 35696286 264259 284892
				הפעופון וווחון ביות		18108362, 33657023, 29331824, 33657109.
						29146499, 264508, 264509, 264905, 264906, Jeanna 264906, 264906, 264909
				•••		35696423, 35695855, 264510, 264511.
						264512, 264910, 264634, 264635, 264637,
	·					264638, 33657402, 264758, 85658542, 264602, 264760, 264781, 264482, 264563.
						264762, 264483, 264764, 264566, 264288, 264768
1908	12763822 (3811, 3812)	1, 3812)			UNCLASSIFIED	264637
	1					

65274572, 22278996, 35696286, 22278998. 22278999, 264259, 60432049, 29331822. 29331825, 60424269, 29331828, 35696052, 284593, 6043336, 21906754, 55811386, 264682, 264684, 264288, 21906765, 21906768, 21906768, 21906768, 21906769, 33657109, 35695855, 264638, 264638, 60432113	264488, 65274572, 56182575, 5016 1000, 22278995, 22278996, 22278997, 22278998, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 204269, 29331822, 29146499, 264905, 264569, 29331828, 29146499, 264905, 264907, 264907, 264909, 264906, 264907, 264908, 264594, 60433438, 264596, 55812038, 33109954, 52646317, 265011, 265017, 26404, 265018, 264607, 26279000, 222790002, 264566, 264486	264259, 264508, 264805, 264806, 254807, 264908, 264511, 264910, 264593, 264758, 264766, 18108370, 264634, 264637, 264486	35696286, 22278996, 22278999, 35696286, 22278996, 22478996, 264907, 264908, 264907, 264908, 264907, 264908, 264909, 264907, 264909, 264910, 264758, 265011, 264601, 264602, 264604, 264768, 264761, 264764, 264268, 264766, 264769, 35695917, 265021, 264629, 18108372, 18108374, 35695855, 264631, 264634, 264635, 264656, 264568, 2646837, 264638, 18108385, 264565, 264568, 264468
	UNCLASSIFIED	UNCLASSIFIED	нотеорох
Contains protein domain (PF00561) - hydrolase alpha/beta hydrolase fold			Contains protein domain (PF00412) - homeobox LIM domain containing proteins
1 AF15181 - (AF151816) CGI-58	95313641 (3815, 3816) Novel Protein sim. GBank gij3986770 (AF109906) - NG22 (Mus musculus)	85514505 (3817, 3818) Novel Protein sim. GBank gi 2224653 dbj BAA20813  - (AB002354) KIAA0356 [Homo sapiens]	94216821 (3819, 3820) gij1351218jspjP47226jTES2_MOUSE - TESTIN 2 (TES2) (CONTAINS: TESTIN 1 (TES1))
1907   95351144 (3813, 3814)   Novel Protein sim. GBank gil4929585[gb AAD34053. protein [Homo sepiens]	1908 95313641 (3815, 3816) N	1909 (85514505 (3817, 3818) N	1910 94216821 (3819, 3820) <sup>1</sup>

			3000000	0000	101/00104 ER10257
1911	1911   91725345 (3821, 3822) Novel Protein sim. GBank	Vovel Protein sim. Gibank Habbaaaalakia Anania 4 (1000653 - (40006530)	ONA mismatch repair profein		29331827, 33656970, 264906, 265007.
		hypothetical protein (Homo sapiens)			264591, 55812038, 87168559, 264448.
					264369, 21906765, 21906768, 265022,
					264691, 264693, 18108365, 55811576,
					264556, 18108385, 18108388
1912	_	95413519 (3823, 3824) Novel Protein slm. GBank gil5689439 dbj BAA83003.1  -	Contains protein domain (PF00098) - UNCLASSIFIED	JNCLASSIFIED	18108397, 56182575, 56181686, 22278994.
!		(AB028974) KIAA1051 protein [Homo sapiens]	Zinc finger, CCHC dass		22278995, 56994075, 22278996, 22278999.
					264259, 29331822, 29331824, 56182181,
					29331825, 66714117, 35696052, 264905,
					264906, 264907, 264908, 52644045,
					56182435, 265007, 265008, 264910, 265009,
					264591, 264596, 65274444, 55811386,
					87168474, 265011, 87168559, 265018,
					265019, 264760, 18108351, 264681, 264369,
					264684, 264288, 264686, 264768, 21906765,
					21908766, 21906767, 21906768, 21906769,
					55811957, 265020, 265021, 265022,
					60170615, 264692, 33657023, 264693,
					18108376 55811576 35696423, 65274791,
					264637 56182323, 83373044, 56526486,
					22279002, 264563, 264566
101		DESTREAM (3828, 3828) Novel Profein eim GBank			56182575, 22278994, 22278995, 56994075,
2	2000000 (0000) 0000)	piis032245imilNP 005665 11p2NF2 - zinc finger protein			22278996, 22278998, 22278999, 29331826.
		(C2H2) homologous to mouse MOK-2			29331827, 265008, 55812038, 265010.
			·		265017, 265018, 265019, 264681, 18108351,
					264683, 264764, 264369, 264288, 264685,
					264686, 264769, 21906765, 21906766,
					21906768, 21906769, 55811957, 265020.
_					265022, 264691, 55811576, 264634, 264635,
					264638, 56182323, 83373044, 18108385
1914	_	83423982 (3827 3828) Novel Protein sim. GBank gil4589604ldbilBAA76824.11 -	Contains protein domain (PF00036) - struct	struct	56182575, 29331824, 35696052, 264908.
<u> </u>		(AB023197) KIAA0980 protein [Homo saplens]	EF hand		264908, 264828, 264909, 264592, 264758,
					87168559, 18108351, 18108354, 264684.
					264686, 33657023, 264893, 264628, 264631.
_			,		264632, 264634, 264635, 264639
1015		95340459 (3829 3830) Novel Protein sim. GBank gil5689415IdbilBA882991.11-		UNCLASSIFIED	264259, 29331824, 29331826, 29331827.
2					264508, 264909, 265009, 265017, 265019,
					264768, 264769, 264689, 264628, 264635,
_					264637, 264639, 83373044, 264353
1916	3 79640761 (3831, 3832)				264693, 264639

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slud	sfruct	oxidase	UNCLASSIFIED	UNCLASSIFIED
1917   87821580 (3833, 3834) Novel Protein sim. GBank gil5689391(dbj BAA82979.1  - (AB028950) KIAA1027 protein [Homo sapiens]	95302795 (3835, 3836) Novel Protein sim. GBank gij5281517[gb]AAD41524.1 AF15483 - (AF154831) PV-1 [Raftus norvegicus]	94143847 (3837, 3838) Novel Protein sim. GBank gij3878584 emb CAB01237  - (Z77667) cDNA EST EMBL: C08125 comes from this gene; cDNA EST EMBL: C09753 comes from this gene	91229953 (3839, 3840) Novel Protein sim. GBank gij1809231 (AC000115) - coded for by human cDNAs R76043 (NID:g850725), R65857 (NID:g838495) and H12868 (NID:g877688) [Homo sapiens]	79555226 (3641, 3842) Novel Protein sim. GBank gi[4580997[gb]AAD24571.1JAF12108 - (AF121081) cAMP inducible 2 protein [Mus musculus]
8782	1918 95300	1919 9414	1920 9122	1921 7955

930	87889128 (3859, 3860)	1930 87889128 (3859, 3860) Novel Protein sim. GBank		phosphatase	35696286, 29331828, 264905, 264907,
		Bilivoszou(spirozsosjubita_moosc - MBL4 TRO IEIN			264601, 265017, 265019, 264605, 264760.
					264764, 264766, 264688, 264769, 265022,
					35696423, 264638, 60432113
1831	87797279 (3861, 3862)	87797279 (3861, 3862) Novel Protein sim. GBank gi 404634 (U01840) -	Contains protein domain (PF00069) - kinase	kinase	264906, 264908, 60432229, 264758, 264764,
		serine/threonine kinase (Mus musculus)	Eukaryotic protein kinase domain		264288, 265020, 264692, 264634, 264637
	15030972 (3863, 3864)				264684, 264691, 264635
	11613668 (3865, 3866)			UNCLASSIFIED	264595
1834	84426360 (3867, 3868)	84426360 (3867, 3868) Novel Protein sim. GBank gif4115748 dbj BAA36494  -		struct	56182575, 56182435, 264510, 264757,
		(AB022023) nonmuscle myosin heavy chain B (Bos taurus)			264758, 55812038, 55811388, 265018,
					55811150, 21906765, 264691, 284631,
	,				264635, 264637
1935	87752511 (3869, 3870)			UNCLASSIFIED	264686, 265011, 264511, 264905, 18108351,   264564 264681 264259 18108370 264568
					264764, 264369, 264595
1936	-	95414338 (3871, 3872) Novel Protein sim. GBank			60432289, 265007, 265010, 265011, 265019,
		~			33657109, 18108374
		receptor-associated protein, 150 kDa subunit			
1937	94847141 (3873, 3874)	94847141 (3873, 3874) Novel Protein sim. GBank gil543187 pir S37771 - ankyrin,	Contains protein domain (PF00023) - kinase	kinase	85658542, 21906767, 35695917, 60170615,
	_	erythrocyte - mouse	Ank repeat		264693, 33657109
1938	_	87403277 (3875, 3876) Novel Protein sim. GBank	Contains protein domain (PF00888) - collagen	collagen	264488, 29146498, 264905, 264559
		giļ454431[gbļAAD22340.1 AC00695 - (AC006955) hvpothetical protein [Arabidoosis thailana]	Cullin family		
1939		91004978 (3877, 3878) Novel Protein sim. GBank gil500858IdbilBAA032101 -		UNCLASSIFIED	65274572, 22278999, 60432289, 56182435.
		(D14168) 50kDa lectin [Bombyx mori]			60433356, 265017, 21906765, 21906766,
					21906768, 55811957, 27486264, 35696423, 60432113, 264688
970	7		10000000	44-144	201700 201700 201710 201700 201700
5			Contains protein domain (Prudoou) - struct Leucine Rich Repeat	struca	264486, 29331622, 264448, 264863, 264289, 265020, 33657023, 264631
1841		94147177 (3881, 3882) Novel Protein sim. GBank gij4206386 (AF060570) - rig-1		UNCLASSIFIED	56994075, 22278998, 264259, 29331824,
		protein [Mus musculus]			29331827, 264905, 265008, 33857084.
					265017, 265018, 264288, 264687, 21908765,
					21906766, 21906767, 265020, 52644150,
					27486264, 83373044, 18108387, 60432113.
	-				22279002, 264565
<del>2</del>				UNCLASSIFIED	264488, 18108398, 29331825, 27486261,
		gi 4927204 gb AAD33049.1 AF13391 - (AF133911) ARL-6  interacting profeln-4 (Mus muscrifus)			264509, 18108370, 18108374, 264482
1843	_	94325298 (3885, 3886) Novel Protein sim. GBank	Contains protein domain (PF00400) - kinase	kinase	22278998, 29331822, 29331827, 35696052,
		gij3122952jspjO15736jTiPD_DICDI - TIPD PROTEIN	WD domain, G-beta repeat		264511, 265009, 264592, 60432229, 265017,
					265018, 265019, 264684, 264692, 33657109,
					65274791, 264636

1844	94232958 (3887, 3888)	1944   94232958 (3887, 3888) Novel Protein sim. GBank gij1799570jdbjjBAA13432] - (7)87871 TIP190 (Bathis noncentrie)		UNCLASSIFIED	65274572, 22278994, 22278995, 22278996,
					22110330, 22210333, 204233, 32043080, 29331824, 29331826, 29331827, 66712502,
					56182435, 264512, 264910, 60170831,
					60432229, 60433356, 33657402, 60433438,
		*			264596, 33109954, 21906754, 87168474,
					87168559, 265017, 265018, 265019,
			•		18108351, 264369, 264686, 264768.
					21906765, 21906768, 21906767, 21906769,
				-	35695917, 265020, 265021, 60170615,
					264692, 33657023, 18108370, 18108374,
					35696423, 35695855, 264634, 60170394,
					264639, 83373044, 18108385, 56526486,
,					87168518, 60432113, 22279000, 264563
C # 6		o/o416/4 (3868, 3890) Novel Protein sim. GBank		UNCLASSIFIED	264488, 22278996, 264510, 264511,
		91492/204 gb AAD33049.1 AF13391 - (AF133911) ARL-6   Interacting protein-4 IMus musculus			18108351, 264683, 264486, 264567
1946		87443990 (3891, 3892) Novel Protein sim. GBank		UNCLASSIFIED	60432289 29331827 35696052 265007
		gip498104 splO27969 AD50_BOVIN - ADRENAL			265008, 60433356, 60433438, 264369.
		MEDULLA 50 KD PROTEIN			56181562, 21906767, 52644150, 264693.
					27486264, 264637, 87168518, 264563
¥.	86438862 (3893, 3894)	86438862 (3893, 3894) Novel Protein sim. GBank		rnapolymerase	22278998, 264905, 264906, 264908, 264909,
		gij3914801 sp O54888 RPA2_RAT - DNA-DIRECTED RNA			264512, 264758, 264762, 264682, 264683,
		POLYMERASE I 135 KD POLYPEPTIDE (RNA			284764, 264288, 264768, 264688, 21906768.
		POLYMERASE I SUBUNIT 2) (RPA135) (RNA			264693, 18108374, 35695855, 264635.
9,0	_	POLYMERASE I 127 KD SUBUNIT)			264637, 264638, 18108385, 22279002
\$				struct	264909, 60170831, 264591, 264594, 235010,
		(AJ243459) proteophosphoglycan [Leishmania major]			265011, 264764, 264369, 264689, 264631,
1040	7640120 /2807 20001				264638
96	_				264369
926	_	of 199331 (3989, 3800) INOVEL Protein Sim. GBank gij3876766jembjCAA93466.1	Contains protein domain (PF00857) - UNCLASSIFIED	UNCLASSIFIED	264488, 264768, 264688, 264689, 264259,
		(209037) predicted using Genetinder; Similarity to E.coli	Isochorismatase family		29331822, 33657109, 29331828, 264103,
		Capacitabdille alegans			264509, 18108370, 35695855, 264510,
					265008, 265009, 33657402, 18108385,
		•			265018, 264563, 55811150, 18108351,
1951		86988253 (3901, 3902) Novel Protein sim GBank oil2626753dhillBAA234241	Control of the state of the sta		264369, 264288, 18108354
		(ABOOR782) euffate transmoder [Ambidomin thelians)	Contains protein domain (Pruggle) - Irransport	ranspor	561825/5, 222/8997, 52645080, 29331824,
		מיישים לא המיישים וישיים לאומסות לא המיישים וישוושנום	Suirate transporter family	`	29331825, 29331827, 55812038, 52646317,
					265018, 265019, 264369, 21906765,
					21908767, 55811957, 265020, 265021,
					33657023, 264693, 35695763, 56182323, 22279002
1952	_	87069775 (3903, 3904) Novel Protein sim. GBank	Contains protein domain (DE00108)	codumpto	DEAEDS DEAEDS DEAEDS DEAEDE DEAEEE
			_	reductase	204591, 204593, 204594, 204595, 204555, 204578, 204577, 204778, 204707
		protein (Homo sapiens)	Snon chain denyarogenase		264556, 264557, 264558, 264565

	20470371 (390	5, 3906)	1953   20470371 (3905, 3906)   Novel Protein sim. GBank gil1168715 sp P31721 C1QB_RAT - COMPLEMENT C1Q. SIBCOMPONENT R CHAIN DECTIDEDE	Contains protein domain (PF00386) - complement C1q domain	complement	264259, 264558	
915	<u> 26025 (390)</u>	7, 3908)	91226025 (3907, 3908) Novel Protein sim. GBank gild240271[db][BAA74914.1] - (AB020698) KIAA0891 protein [Homo saplens]	Contains protein domain (PF00443) - ubiquitin Ubiquitin carboxyl-terminal hydrolase family 2	ubiquitin	264488, 263994, 18108394, 356962286, 22278998, 29331822, 66714117, 29331826, 29331827, 264508, 264509, 264905, 264906, 264907, 264908, 264909, 264905, 264906, 265007, 265008, 264910, 265009, 60170831, 265019, 18108351, 264685, 264766, 264769, 35695017, 265021, 264691, 33657023, 264692, 33657023, 264692, 33657023, 264636, 264585, 264536, 264536, 264536, 264536, 264536, 264536, 264536, 264536, 264536, 264536, 264556, 264563, 264563, 264556, 264564, 264565, 264563, 264563, 264563, 264564, 264566, 264666, 264	
ξς -	95308310 (3909, 3910)	9, 3910)			UNCLASSIFIED	56182575, 56181686, 22278994, 22278999, 264259, 264259, 264259, 29331822, 56182181, 29331824, 29331824, 29331825, 29331825, 29331827, 35696052, 264508, 29331830, 265009, 264509, 264509, 264509, 264409, 26440, 265017, 265018, 265019, 264448, 26461, 21906765, 21906766, 21906766, 21906766, 21906766, 265021, 52645129, 33657109, 27486264, 33657149, 35695763, 60431528, 18108374, 264639, 83373044	
S6	092121 (391	1, 3912]	95092121 (3911, 3912) Novel Protein sim. GBank gi 1865821 dbj BAA13407  - (D87469) Similar to D.melanogaster cadherin-related tumor suppressor [Homo sapiens]	Cadherin domain (PF00028) - cadherin	cadherin	264488, 56182575, 22278996, 22278999, 264259, 29331827, 35698052, 29331824, 29331825, 29331827, 35698052, 29331828, 264508, 264906, 264909, 264910, 26490, 264910, 264910, 264768, 56181562, 264689, 19005768, 21906769, 265022, 264691, 265028, 18108374, 55810764, 55811576, 264628, 18108374, 55810764, 55811576, 264639, 60170394, 56182323, 83373044, 18108385, 22279000, 22279002, 264563, 264564	

1957	94326510 (3913, 3814)			UNCLASSIFIED	52646842, 56182575, 22278997, 22278998, 22278999, 29331824, 66714117, 29331827, 29346498, 264593, 33657402, 33109954, 87168474, 265018, 264448, 264369, 264288, 264766, 21906766, 21908767, 21906768, 21906769, 265021, 264692, 65274620, 27486264, 33657349, 27486265, 35695855, 22279002, 264482
	95313902 (3915, 3916)	95313902 (3915, 3916) Novel Protein sim. GBank gil4240227 dbj BAA74892.1  - (AB020676) KIAA0869 protein [Homo sapiens]		UNCLASSIFIED	22278999, 264092, 264094, 264259, 60432049, 29331824, 56162181, 66714117, 264107, 264109, 264909, 264511, 60170831, 60432229, 21906754, 265010, 21906769, 35695917, 265022, 65274620, 263967, 263976, 35696423, 264631, 264632, 264634, 264635, 18108385, 22279000, 22279002
		85701470 (3917, 3918) Novel Protein sim. GBank gij2281983jembjCAB10860j - (298056) hyypothetical protein (Schizosaccharomyces pombe]		ubiquitin	<u>2</u> 64593, 265019
		80308608 (3919, 3920) Novel Protein sim. GBank gi 2274851 db  BAA21515  - (D84159) 3-7 gene product [Homo sapiens]		struct	264905, 264906, 264907, 264908, 264909, 265006, 265007, 264910, 264595, 265017, 264604, 265018, 18108351, 264764, 264369, 264766, 264768, 21906765, 18108369, 264639, 1810839, 264635, 264636, 264637, 264638, 264637, 264637, 264636, 264636, 264637, 264636, 264636, 264637, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264636, 264637, 264636, 264636, 264637, 264636, 264636, 264636, 264637, 264636, 264636, 264636, 264637, 264636, 264636, 264637, 264636, 264636, 264637, 264636, 264636, 264637, 264636, 264636, 264637, 264636, 264636, 264637, 264636, 264636, 264637, 264636, 264636, 264637, 264636, 264636, 264637, 264636, 264636, 264636, 264636, 264636, 264636, 264637, 264636, 264636, 264637, 264636, 264646, 264666,
_	16292607 (3921, 3922)				264635
795	91008385 (3923, 3924)			UNCLASSIFIED	65274572, 264592, 264593, 265019, 264691
	90936017 (3925, 3926)			ерћ	65274572, 18108398, 35696286, 29331825, 60432289, 29331827, 264828, 265006, 285009, 60433356, 60433438, 21906754, 265020, 265021, 33657023, 33657109, 27486265, 35695855, 264555
	94317605 (3927, 3928)	9431 7609 (3927, 3928) Novel Protein sim. GBank gi[3262638 emb CAB45757.1  - (AL080169) hypothetical protein [Homo sapiens]		cadherin	264488, 264092, 264259, 264509, 264905, 264906, 264907, 264908, 264909, 264510, 264908, 264909, 264510, 264911, 265007, 265009, 264910, 264592, 264593, 264594, 264595, 264788, 264600, 264603, 284604, 284805, 284760, 264768, 264768, 264768, 264769, 264769, 264769, 264691, 264692, 264693, 18108370, 264629, 264634, 264639, 264637, 264638, 18108372, 264638, 18108382, 264433, 264564, 264565, 264566, 18488, 284567, 264868, 284565, 264566, 264868, 284565, 264566, 264868, 284567, 284868, 284565, 284566, 28466, 28466, 28466, 28466, 28466, 28466, 28466, 28466, 28466, 28466, 28466, 28466, 28466, 28466, 28466, 28466, 28466, 28466, 28466, 2846
1865	94317445 (3929, 3930)	94317445 (3928, 3930) Novel Protein sim. GBank gil4107017 dbj BAA36294  -  (AB001773) PEM-6 [Ciona savignyi]	Contains protein domain (PF01428) - ubiquitin AN1-like Zinc finger	ubiquilin	264488, 264510, 264760, 264768, 264486

22276999, 264092, 264259, 29331826, 29331828, 29146498, 264595, 265011, 264448, 18108354, 264288, 264684, 264766, 264685, 264688, 265022, 264691, 264692, 18108370, 18108377, 264555, 18108381, 18108395, 264486, 264567	29331824, 265010, 265017, 264288, 21906764, 263981, 56526488				264486, 264487, 222,10930, 264428, 264486, 26432249, 260432249, 26331827, 35696052, 264508, 264905, 264509, 264509, 264509, 264509, 264500, 264906, 264906, 264501, 264511, 264512, 264908, 26450, 264500, 264604, 264605, 264600, 18108351, 264764, 264764, 264288, 264766, 264768, 264764, 264288, 264769, 21906765, 33657023, 264692, 18108370, 264629, 35696423, 264635, 264636, 264631, 264555, 264631, 264563, 264636, 264564, 264654, 264636, 264638, 264637, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264663, 264486			IED 264682, 264764, 264563
glycoprotein	esterase	oncogene	UNCLASSIFIED	UNCLASSIFIED	collagen		UNCLASSIFIED	UNCLASSIFIED
Contains protein domain (PF0 1027) - glycoprotein Uncharacterized protein family	Contains protein domain (PF00388) - Phosphatidylinositol-specific phosphotipase C, X domain	Contains protein domain (PF00008) - oncogene EGF-like domain				Contains protein domain (PF01462). Leucine rich repeat N-terminal domain		,
Contains protein domain (PF01 gi 4929707 gb AAD34114.1 AF15187 - (AF151877) CGI-119 Uncharacterized protein family protein family protein [Homo saplens]	87396123 (3933, 3934) Novel Protein sim. GBank gil2957270 (AF044576) - phospholipase C PLC210 [Caenomabditis elegans]	88095641 (3935, 3936) Novel Protein sim. GBank gi[2564953 (AF030001) - unknown [Mus musculus]	Novel Protein sim. GBank gi [Nephila clavipes]	80596049 (3939, 3940) Novel Protein sim. GBank gil4050087 (AF109907) - S164 [Homo saplens]	94843914 (3941, 3942) Novel Protein sim. GBank gij134206jsp P09593 SANT_PLAFV - S-ANTIGEN PROTEIN PRECURSOR	87645444 (3943, 3944) Novel Protein sim. GBank gij4519623[dbj BAA75671.1] - (AB017616) homologous to the yeast YGR163 gene [Musmusculus]		80396629 (3947, 3948) Novel Protein sim. GBank gij3309543 (AF036382) - MLL
94192058 (3931, 3932) N	87396123 (3933, 3934) P	86095641 (3935, 3936) (	84328529 (3937, 3938)	80596049 (3939, 3940)	94843914 (3941, 3942)		86395533 (3945, 3946)	_
996	1967	1968	1969	1970	1871	1972	1973	1974

UNCLASSIFIED   264488, 66714117, 29331828, 29331828,	UNCLASSIFIED 284259, 35696052, 265018, 265020, 265021, 33657109, 56528486	hотеоbох 264908, 264596, 265021, 264566	transcriptfactor 60170831, 264566	UNCLASSIFIED 29331626, 29146498, 264905, 264907, 265007, 265009, 265010, 285018, 264686, 18108359, 21906768, 35695917, 265020, 60170815, 264693, 18108368, 18108370, 264635, 264556, 264558, 18108384, 22279000, 264545	ubiquitin 264489	UNCLASSIFIED 22278995, 22278997, 22278999, 264259, 29331825, 29331828, 29146498, 29146499, 264107, 264908, 264910, 264595, 21906754, 265017, 265018, 265019, 264448, 264288, 21906767, 33657023, 27486264, 18108370, 18108374, 18108374, 18108376, 264631, 264631, 264631, 264487, 264631, 264487, 264631, 264487, 264631, 264631, 264487, 264631, 264631, 264487, 264631, 264631, 264487, 264631, 264487, 264631, 264	6574572, 29331825, 35696052, 33656970, 264909, 265008, 55811386, 264760, 284688, 26486, 26486584	
	,	4	in .					Contains protein domain (PF00086) - Iranscriptfactor Zinc finger, C2H2 type
(0		94852664 (3953, 3954) Novel Protein sim. GBank gil2499526 sp QQ7782 NASU_RAT - SODIUM/SULFATE COTRANSPORTER (NA(+)/SULFATE COTRANSPORTER)	87447645 (3955, 3956) Novel Protein sim. GBank gi]103421 pirj A33471 - transcription factor NTF1 - fruit fly (Drosophila melanogaster) (fragment)	87627709 (3957, 3958) Novel Protein sim. GBank gil2244815 emb CAB10238.1  - (297336) hypothetical protein [Arabidopsis thaliana]	86577059 (3959, 3860) Novel Protein sim. GBank gil4759290[ref]NP_004642.1[pUSP1 - Ubiquitin carboxyl- terminal hydrolase, X-linked	87606974 (3961, 3962) Novet Protein sim. GBank gil4406693 gb AAD20060  - (AF131849) Unknown [Homo sapiens]	90995367 (3963, 3964) Novel Protein sim. GBank gij5689523 dbj BAA83045.1  - (AB029016) KIAA1093 protein [Homo sapiens]	95098668 (3965, 3966) Novel Protein sim. GBank gij3417297 (AC002310) - Unknown gene product [Homo sapiens]
1975   94316479 (3949, 3950)		1977   94852664 (3953, 3954)						1983   95098668 (3965, 3966)

94122108 (3975, 3974) Novel Protein sim. CBank [Mesocricetus auratus] 94122108 (3975, 3976) Interacting protein kinase PKM [Mesocricetus auratus] 941225225 (3977, 3976) Novel Protein sim. GBank gil2801701 (AF042379) - spindle pole body protein spc97 homolog GCP2 [Homo sapiens] 85699888 (3979, 3980) Novel Protein sim. GBank gil5701727[db][BAA83074.1] - (AB024729) atpha-1.3-D-mannoside beta-1.4-N-acetylglucosaminyltransferase IV-homologue [Homo sapiens] 85699888 (3981, 3982) Novel Protein sim. GBank gil4240287[db][BAA74922.1] - Con (AB020706) KIAA0899 protein [Homo sapiens]	Contains protein domain (PF00069) -  Eukaryotic protein kinase domain  UNCLASSIF  tubulin  tubulin  Adaptin N terminal region	OH CONTRACTOR OF THE CONTRACTO	264488, 264629 22278999, 29331830, 265007, 265018, 21906768, 33657023, 264692, 264693, 21906768, 33657023, 264692, 264693, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264906, 264639, 264630, 264639, 264638, 264639, 264630, 264639, 264636, 264630, 264764, 18108381 18108394, 56182575, 22278994, 35696286, 56994075, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22278997, 22378997, 22378997, 22378997, 22331828, 263331824, 29331824, 29331824, 29331828,
95317232 (3983, 3984) Novel Protein sim. GBank gil2246532 (U93872) - ORF 73, Go contains large complex repeat CR 73 [Kaposi's sarcomaassociated herpesvirus] 80054763 (3985, 3986) Novel Protein sim. GBank gil2565091 (U80761) - CTG26	Contains protein domain (PF00096) - struct	sind 	29331824, 29331825, 60432289, 29331828, 264508, 264906, 264907, 264908, 56182435, 264508, 264906, 264906, 264907, 265008, 56182435, 264510, 265007, 21906764, 33109954, 31109351, 264507, 265018, 265019, 264762, 18108351, 264769, 264683, 264769, 21906768, 21906768, 21906769, 55811957, 265020, 60431528, 263974, 18108379, 35695855, 264555, 264557, 264639, 83373044, 18108384, 87168518, 60432113, 22279000, 22279002, 264564, 264468, 264766, 264608, 264768, 264768, 264768, 264768, 264768, 264768, 264768, 264568, 264568, 264567, 2645767, 264767, 2645767, 2645767, 2645767, 2645767, 2645767, 2645767, 2645767, 2645767, 2645767, 2645767, 2645767, 2645767, 2645767, 26457

264488, 22278997, 22278998, 22278999, 29331822, 29331824, 29331825, 29331825, 29331826, 29331825, 29331826, 29331827, 29331828, 264905, 264907, 264908, 265008, 60432229, 33657402, 60433356, 264757, 60433438, 264758, 21906764, 264394, 264369, 264686, 264768, 265020, 264690, 18108362, 264693, 264562, 18108370, 264635, 264555, 26456486, 22279000, 22279002, 264564	264488, 18108396, 22278994, 56994075, 22278996, 35696286, 22278997, 22278999, 224278996, 35696286, 22278997, 22278999, 224278996, 3569628, 22278997, 22278999, 2242289, 29341827, 35696052, 29146499, 284905, 264907, 66712502, 56182435, 265008, 265007, 265008, 265009, 265010, 265011, 265017, 265018, 265019, 265010, 265011, 265017, 265018, 265019, 264761, 264762, 264689, 21906765, 21906767, 29148627, 21906765, 21906766, 21906767, 29148627, 21906768, 21906769, 2914874, 265020, 265021, 264690, 18108377, 35698423, 55811576, 65274791, 3569855, 264634, 264558, 264556, 264557, 264558, 264554, 264557, 264557, 284558, 264554, 264557, 264557, 264558, 2645113, 264557, 264557, 264557, 264558, 2645113, 264557, 264567, 26457, 264567, 26457, 26457, 26457, 26457, 26457, 26457, 26457, 26457,	264564	264908, 264909, 264592, 264593	264113, 264685, 264555, 264567
,	UNCLASSIFIED	UNCLASSIFIED	phosphatase	UNCLASSIFIED
Contains protein domain (PF00856) - mapolymerase SET domain			Contains protein domain (PF00023) - phosphatase Ank repeat	
1994   94329114 (3987, 3988) Novel Protein sim. GBank gi 5630077 gbpAAD45622.1AC00601 - (AC006017) similar to ALR; similar to AAC51735 (PID:g2358287) [Homo sapiens]	1895 95414353 (3989, 3990) Novel Protein sim. GBank gil4827040 ref NP_005110.1 pTRAP - thyrold hormone receptor-associated protein, 150 kDa subunit	80254186 (3991, 3992) Novel Protein sim. GBank gij791146jembjCAA60020j - (X86028) extensin-like protein [Vigna unguiculata]	87028423 (3993, 3994) Novel Protein sim. GBank gilz642034 (AF034547) - protein phosphatase M130 myosin binding subunit [Ovis aries]	85262704 (3995, 3996) Novel Protein sim. GBank gil4589634 dbj BAA76839.1  -   (AB023212) KIAA0995 protein (Homo sapiens)
1994 9432	1995 95414			1998 8526

			264907, 285008, 284910, 284603, 264892, 264629, 18108374, 264556, 264557	(IED 264905, 264908, 264908, 264910, 264596, 265017, 18108351, 264692, 264629, 264634, 264565
NCLASSIFIE	UNCLASSIFIED	UNCLASSIFIED	isomerase	UNCLASSIFIED
Contains protein domain (PF00515) - UNCLASSIFIED TPR Domain			Contains protein domain (PF00085) - isomerase Thioredoxin	
1/AF07244 - (AF072441) i cabin 1 [Homo sapiens]	Novel Protein sim. GBank gij 1723232 sp Q10155 YATA_SCHPO - HYPOTHETICAL 90.6 KD PROTEIN C1D4.10 IN CHROMOSOME I	95072534 (4001, 4002) Novel Protein sim. GBank gij107560jpirj B38637 - Ras inhibitor (clone JC265) - human (fragment)	80236368 (4003, 4004) Novel Protein sim, GBank Golfabar gil729433lsplP38657 ER60_BOVIN - PROBABLE PROTEIN Thioredoxin DISULFIDE ISOMERASE ER-60 PRECURSOR (ERP60) (58 KD MICROSOMAL PROTEIN) (P58)	80074449 (4005, 4006) Novel Protein sim. GBank gij85388jpir  A27040 - neurofilament triplet M protein - chicken (fragment)
1899   94324903 (3997, 3998)   Novel Protein sim. GBank gij5225312 gb AAD40846. calcineurin binding protein	95413705 (3999, 4000) Novel Protein sim. GBank gil1723232lsp Q10155 YA 90,6 KD PROTEIN C104.	15072534 (4001, 4002) h	80236368 (4003, 4004)	80074449 (4005, 4006)
1899	2000	2001	2002	2003

2004	95317318 (4007, 4008)	2004   95317318 (4007, 4008) Novel Protein slm. GBank gl 4884249 emb CAB43230.1  - (AL049996) hypothetical protein [Homo sapiens]	Contains protein domain (PF00076) - dna_ma_bind RNA recognition motif. (a.k.a. RRM,	dna_ma_bind	52645156, 52646842, 52646365, 56182575, 22278996,
			RBD, or RNP domain)		35696286, 22278997, 22278998, 22278999, 60432049, 264259, 52645080, 29331824, 29331826, 29331827, 35698052, 29331828,
					33656970, 29331830, 264908, 264592, 60433356, 33657402, 52646317, 21906754
					33657084, 52644296, 87168474, 87168559,
					265017, 265018, 265019, 284763, 264683, 264288, 62644230, 24006768, 24006768
					21906767, 21906768, 21906769, 35695917,
					265020, 265021, 265022, 52644150,
					33657023, 52645129, 33657109, 33657182,
					2/486261, 2/486262, 3355/349, 2/486265, 35695763, 18108374, 18108376, 18108377
					35696423, 35695855, 264631, 52644332,
					264558, 18108385, 56526486, 87168518,
2005	87400864 (4009, 4010)	87400864 (4009, 4010) Novel Protein sim. GBank gij3879501 jembjCAA87795 j		ubiquitin	264488, 264908
		(247812) similar to ubiquitin carboxyl-terminal hydrolase;			*
		CUNA EST EMBL: U33366 comes from this gene; CDNA			
		EST EMBL: U33953 comes from this gene; cUNA EST EMBL: 033822 comes from this gene: cDNA EST			
[8		EMBL.D34547 comes from this ge			
9007	(2104,1104) /1116668	Novel Protein sim. GBank giļ4106673jemb CAA22613  -	Contains protein domain (PF01702) - UNCLASSIFIED	UNCLASSIFIED	56182575, 56181686, 22278995, 22278996,
		(AL035064) queuine trna-ribosyltransferase [Schiposaccharomycos nombol	Queuine tRNA-ribosyltransferase		22278997, 22278998, 22278999, 264259,
_					56182181, 60424269, 66714117, 35696052,
		۵			264906, 66712502, 264909, 264510. 6043336
					265019 264682 264448 264288 264768
					29148627, 21906769, 29148784, 35695917,
					60170615, 264691, 33657023, 65274620,
					33657109, 55810764, 55811576, 35695855,
2007	94325556 (4013, 4014)	94325556 (4013, 4014) Novel Protein sim. GBank gil2662161 dbj BAA237121 -		UNCLASSIFIED	8/168518, 60432113, 264563, 264482 264488, 263994, 35696052, 264508, 264905
		(AB007900) HH0452 cDNA clone for KIAA0440 has a 438-			264906, 264907, 264908, 264909, 265007.
		be insertion at position 1711 of the sequence of KIAA0440.			264910, 264592, 264595, 264758, 265011,
		[Homo saptens]		•	264760, 264762, 264764, 264766, 264685,
_					264767, 264768, 264769, 55811957,
					35695917, 265020, 264691, 264693, 264628,
					264629, 65274791, 35695855, 264631,
	_				284632, 264634, 264635, 284637, 284638, 264839, 284568
2008			Contains protein domain (PF00046) - homeobox	нотеорох	264909, 264768, 35695855
		I(TO/SOU) nomeodomain protein (Mus musculus)	Homeobox domain		

22278999, 264259, 264910, 264591, 265017, 264681, 264683, 21906768, 264691, 33657182, 33657349, 264631, 87168518, 264404, 22279002, 264563	52644507, 52645156, 52646365, 52646842, 18108397, 65274572, 22278994, 55994075, 35696286, 22278994, 56994075, 35696286, 22278994, 26931827, 22278999, 22278999, 264559, 264559, 29331822, 2645508, 29331824, 29331825, 29331822, 2656508, 265009, 6043229, 60433356, 21906754, 52646317, 33109954, 52644296, 87168474, 87168559, 265017, 265018, 265019, 264681, 264685, 264697, 5264150, 264691, 264692, 35695917, 265020, 52644150, 264691, 264692, 3569563, 36557023, 263967, 52645129, 35695763, 18108376, 3569423, 65274791, 35695855, 2644332, 601470394, 18108385, 87168518, 22279002, 284564, 264565, 264566, 264567	56182575, 56994075, 22278999, 264259, 29331824, 29331824, 29331824, 29331824, 29331824, 29331824, 29331824, 2959065, 2659066, 265907, 265007, 265010, 264010, 264369, 264268, 264685, 264468, 264687, 21906767, 21906768, 21906768, 21906769, 265020, 265022, 264691, 33657023, 65274620, 33657109, 264629, 264557, 264559, 83373044, 87168518, 60432113, 22279902	264259, 29331822, 29331824, 29331825, 2946498, 87168559, 265019, 264682, 264288, 264686, 21906764, 265020, 265022, 33657023, 264693, 33657109, 55811576, 264632, 264558, 56182323, 264639, 18108388, 18108388, 264567	18108398, 264908, 265007, 265010, 265018, 265019, 264689, 21908767, 265020, 284692	264488, 29331826, 264907, 264636, 264555, 264639, 264558
UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED	dna_ma_bind	UNCLASSIFIED
	Contains protein domain (PF00057) - eph Low-density lipoprotein receptor domain class A		Contains protein domain (PF00409) - UNCLASSIFIED Kinesin light chain repeat	Contains protein domain (PF00096) -   dna_ma_bind   Zinc finger, C2H2 type	
2009   85749240 (4017, 4018) Novel Protein sim. GBank gil3882305 dbj BAA34512.1  - (AB018335) KIAA0792 protein [Homo sapiens]	85422458 (4019, 4020) Novel Protein sim. GBank gil5262629JemblCAB45753.1	94328149 (4021, 4022) Novel Protein sim. GBank gil3347953 (AF076183) - cytosolic sorting protein PACS-1a [Rattus norvegicus]	87772137 (4023, 4024) Novel Protein sim. GBank gij1086678 (U41020) - coded for by C. elegans cDNA yk100g4.5; coded for by C. elegans cDNA yk100g4.3; weakly similar to human SREBP-2 basic-helix-loop-helix-leucine zipper (ranscription factor [Caenorhabdilis elegans]	94843842 (4025, 4026) Novel Protein sim. GBank gil4507985 ref NP_003427.1 pZNFf - zinc finger protein 135 Zinc finger, C2H2 type (done pHZ-17)	87347940 (4027, 4028) Novel Protein sim. GBank gij127720lspjP20938jMYPO_HETFR - MYELIN PO PROTEIN PRECURSOR
85749240 (4017, 4018)	85422458 (4019, 4020)				
2009	2010	2011	2012	2013	2014

15	88094922 (4029, 4030)	2015   88094922 (4029, 4030) Novel Protein sim. GBank gil81286 pir  522697 - extensin -		UNCLASSIFIED	56182575, 35696286, 264259, 35696052.
		Volvox carlen (fragment)			264504, 264504, 264584, 264588, 264689, 264628, 264689, 264638, 36695855, 264639, 264563, 264564
2016	85298641 (4031, 4032)	85298641 (4031, 4032) Novel Protein sim. GBank gil285046 pir  S26413 - 1-complex protein Tcp-10 - mouse		stnd	264102, 264508, 264110, 265009, 33109954, 21905768, 265021, 33657109, 27486262, 263972, 18108374, 263978, 264555, 264564
1	79464293 (4033, 4034)			UNCLASSIFIED	284685, 284836
2018	79637067 (4035, 4036)	79637067 (4035, 4036) Novel Protein sim. GBank gi 124735 sp P18175 INVO_PIG-		۲	264693
2019	87787900 (4037, 4038)	87787900 (4037, 4038) Novel Protein sim. GBank gil2143910 pir  568216 - phosphatase-1 giycogen-binding (GL)-chain - rat		phosphatase	264107, 264110, 264112, 265017, 263976
2020		94674476 (4039, 4040) Novel Protein sim. GBank gil2078483 (U43200) - antifreeze glycopeptide AFGP polyprotein precursor (Boreogadus saida)		UNCLASSIFIED	264259, 264508, 264591, 265018, 264682, 264288, 264688, 22278002
2021	86718818 (4041, 4042)	86718818 (4041, 4042) Novel Protein sim. GBank gijs85084[spj007803]EFGM_RAT - ELONGATION FACTOR G, MITOCHONDRIAL PRECURSOR (MEF-G)			56994075, 264593, 33109954, 21906754, 21906768, 33657023, 33857109, 27486261, 87168518
2022	95295665 (4043, 4044)	95295665 (4043, 4044) Novel Protein sim. GBank gil4218005 (AC006135) - putative vicilin storage protein (globulin-like) [Arabidopsis thaliana]			264757, 264767, 60170515, 18108385
923	87722976 (4045, 4046)	2023 87722976 (4045, 4046) Novel Protein sim. GBank gij5410230jgb AAD42992.1 AF07334 - (AF073344) ubiquitin Ubiquitin carboxyl·lerminal specific protease 3 [Homo sapiens]	Contains prolein domain (PF00442) - ublquitin Ublquitin carboxyl-terminal hydrolases family 2	ubiquitin	18108394, 22278999, 264259, 264905, 264906, 264908, 264595, 264762, 264769, 264634, 264636, 87168518, 60432113, 22279000, 264482, 264565
2024	87896443 (4047, 4048)				60433438, 265017, 264688, 264692, 264693, 264636
025	87858863 (4049, 4050	2025 87858863 (4049, 4050) Novel Protein sim. GBank gil4755188[gb]AAD28055.1 AC00701 - (AC007018) unknown protein (Arabidoosis thaliana)	Contains protein domain (PF00637) - UNCLASSIFIED 7-fold repeat in Clathrin and VPS	UNCLASSIFIED	22278997, 264509, 264906, 264909. 55812038, 265017, 265021, 265022. 60170615, 264556
026	94122114 (4051, 4052	2026   94122114 (4051, 4052) Novel Protein sim. GBank gil1655899 emb CAA69032  - (Y07752) pherophorin-S [Volvox carteri]		UNCLASSIFIED	56994075, 60432049, 264508, 66712502, 264112, 60170831, 87168559, 264288, 264689, 21906766, 33657109, 18108385, 60432113, 22279000, 22279002, 264564, 264566, 264567
2027	80249001 (4053, 4054)	0		UNCLASSIFIED	263978, 264634, 264486

52644507, 52645156, 52646842, 56182575, 56994075, 35696286, 22278997, 22278998, 22278998, 22278999, 22278999, 22278999, 22278999, 22278999, 22378997, 22278998, 22378999, 29331824, 56714117, 29331825, 29331826, 29331826, 29331826, 29331826, 29331826, 29331826, 264968, 264967, 264969, 26496745, 265019, 265021, 26502, 25644150, 33657023, 35695763, 246529, 55811576, 35696432, 25695855, 264635, 264636, 2726019, 265019, 265021, 26502, 266036, 27486264, 27486265, 35695855, 264635, 264636, 27270070, 26262	265009, 264595, 85658542, 264555, 264556, 264555, 264556, 264557, 264558, 264559, 81171044	264259, 264326, 22278999, 264259, 26317044 18108394, 56994075, 22278997, 22278999, 264259, 29331824, 29331825, 66714117, 60432289, 29331824, 29331825, 66712502, 264828, 265009, 265019, 265619, 264689, 21906769, 265022, 264682, 264689, 21906769, 265022, 264692, 264693, 264658, 18108370, 18108374, 264634, 264636, 264556, 18108385, 87168518, 22279002, 264565	264591, 55811957, 18108365, 264557,	65274572, 35696286, 29331824, 264908, 265274572, 35696286, 29331824, 264908, 264769, 21906766, 21906767, 29148627, 264628, 35696423, 264634, 264558, 18108381, 60170394, 264559, 83373044, 18108383, 564482, 264684	56181562, 264828, 284632, 264555, 264556	22278996, 22278999, 264907, 29331830, 265008, 285018, 284681, 264682, 264684, 21906767, 21906768, 21906769, 33657109, 83373044, 56526486
ubiquitin	UNCLASSIFIED	transcriptfactor		hydrolase	synthase	dna_ma_bind
Contains protein domain (PF00632) - ubiquitin HECT-domain (ubiquitin- transferase).	Contains protein domain (PF00621) - UNCLASSIFIED RhoGEF domain	Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type	-	Contains protein domain (PF00884) - hydrolase Sulfatase		Contains protein domain (PF00076) - dna_ma_bind RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)
gil 1330345 (U58755) - coded for b1.5; coded for by C. elegans or by C. elegans cDNA yk46e8.5; coded for by C. coded for by C. coded for by C. coded for by C. elegans cDNA	95362032 (4057, 4058) Novel Protein sim. GBank gij3599940 (AF017368) - faciogenital dysplasia protein 2 [Mus musculus]	Novel Protein sim. GBank gl 5630080 gb AAD45825.1{AC00489 - (AC004890) similar to HUB1; similar to BAA24380 (PID:g2789430) [Homo sapiens]		gij5689491[dbjjBAA83029.1] - stein [Homo sapiens]	37_RAT - ALPHA-N- IDE ALPHA-2,6- TGGALNACIII) (STY)	Ę
94699884 (4055, 4056)	95362032 (4057, 4058)	91213734 (4059, 4060)	80245281 (4061, 4062)	91232607 (4063, 4064)	95000809 (4065, 4066)	91232529 (4067, 4068)
2028	-				2033	\$607

2026	93553454 (40E0 4070)				764360 364606 366033 66636406 364667
2038	87115833 (4071 4072)				20231827 20231828 264682 264360
					29148627, 60432113
2037	94324833 (4073, 4074)	94324833 (4073, 4074) Novel Protein slm. GBank gil2734081 (AF000195) - similar		UNCLASSIFIED	65274572, 22278995, 22278996, 56994075.
		to oxysterol-binding proteins [Caenorhabditis elegans]			35696286, 22278997, 22278998, 22278999.
					264259, 29331824, 60432289, 29331826,
					29331828, 35696052, 264907, 29331830.
					66712502, 56182435, 265008, 265009,
		•			60170831, 264594, 55812038, 33109954,
					21906754, 87168559, 265017, 265018,
					265019, 264762, 264369, 264288, 21906765,
					21906767, 21906768, 21906769, 55811957.
					35695917, 265020, 265021, 265022,
				,	52644150, 33657023, 33657109, 33657182,
					35695763, 35695855, 284632, 264634,
					264636, 56182323, 83373044, 60432113,
					22279000, 22279002, 264563
2038	95422384 (4075, 4076)	95422384 (4075, 4076) Novel Protein sim. GBank gij3880625jemb[CAB07858] -	Contains protein domain (PF01412) - UNCLASSIFIED	UNCLASSIFIED	22278995, 22278996, 56994075, 264259,
		(293785) predicted using Genefinder; similar to RNA	Putative GTP-ase activating protein		29331824, 35696052, 264905, 264906,
		recognition motif. (aka RRM, RBD, or RNP domain); cDNA	for Arf		52644045, 265007, 265009, 87168559,
		EST EMBL: T01682 comes from this gene; cDNA EST			265017, 18108351, 264448, 264369, 264766.
		EMBL:M75823 comes from this gene; cDNA EST			264767, 264686, 18108358, 21906765,
		EMBL:D27559 comes from this ge			21906769, 52644150, 33657023, 264692,
				_	18108362, 33657109, 27486262, 18108370,
				•	18108374, 18108379, 35696423, 65274791,
					264632, 264636, 18108383, 83373044,
	=				18108385, 87168518, 22279000, 22279002,
					264563, 264564, 264566
5039	85514626 (4077, 4078)	2039   85514626 (4077, 4078)   Novel Protein sim. GBank gi 2224653 dbj BAA20813  -	Contains protein domain (PF00097) - UNCLASSIFIED	UNCLASSIFIED	22278997, 264259, 29331822, 264905,
		(AB002354) KIAA0356 [Homo sapiens]	Zinc finger, C3HC4 type (RING		264906, 264907, 264908, 264909, 264510,
			(finger)		265009, 264910, 264593, 264758, 265011,
	-	_			265018, 264762, 264288, 264766, 264768,
					264769, 21906766, 33657023, 264692,
					264693, 33657109, 35696423, 264631,
					264632, 264634, 264635, 264636, 264637,
	_				264639, 87168518, 264486
2040	95308417 (4079, 4080)			UNCLASSIFIED	264592
2041	95071736 (4081, 4082)	95071736 (4081, 4082) Novel Protein sim. GBank		mapolymerase	264488, 22278998, 35696052, 264905,
		gij2500625 spjP70700 RPA2_MOUSE - DNA-DIRECTED			264907, 264908, 264910, 265018, 264605.
		RNA POLYMERASE I 135 KD POLYPEPTIDE (RNA			265019, 18108351, 264766, 264769,
		POLYMERASE I SUBUNIT 2) (RPA135)			21906766, 265021, 285022, 264692,
					33657109, 264628, 264629, 35696423,
					35695855, 264637, 264638, 264563, 264564,
					264565, 264567

INCLASSIFIED 60424179, 35696286, 264259, 29331826, 35696052, 29331828, 264508, 264509, 264509, 264501, 264509, 264509, 264510, 264511, 265009, 264910, 264591, 60433356, 264595, 265017, 265019, 264681, 264764, 264369, 264765, 264684, 264788, 284786, 264686, 52644229, 264789, 21808765, 3695917, 264535, 52644150, 264691, 264692, 18108334, 35696423, 65274791, 35695855, 264555, 264556, 22279000, 22279002, 2247902, 264583, 264586, 264566, 264566, 264566		siruct 264093, 29331827, 264905, 66712502, 264592, 264689, 21906765, 21906769, 265020, 264692, 264482, 264568	264692 264259, 264906, 264683, 22279002	22278999, 29147620, 29331824, 29146498, 264508, 265007, 265008, 265019, 264605, 264681, 29148627, 29148629, 265021, 33657023, 18108365, 33657109, 33657182, 18108377, 264556, 264638, 264559, 18108388	UNCLASSIFIED 264909	UNCLASSIFIED 264488, 264259, 29331824, 29331828, 35696052, 284908, 264907, 264908, 264909, 264903, 264763, 21906767, 21906768, 264629, 264634, 264637, 22279002, 264564, 264565, 264566, 264566
Contains protein domain (PF00568) - UNCLASSIFIED WH1 domain	Contains protein domain (PF00122) - transport E1-E2 ATPase	jis				
2042   95307447 (4083, 4084)   Novel Protein sim. GBank gil4406590]gb AAD20040  - (AF131766) Similar to Ena-VASP like protein [Homo sapiens]	94328076 (4085, 4086) Novel Protein slm. GBank gi 5052554 gb AAD38607.1 AF14563 - (AF145632) BcDNA.GH06032 [Drosophila melanogaster]	87106927 (4087, 4088) Novel Protein sim. GBank gi[2246532 (U93872) - ORF 73, contains large complex repeat CR 73 [Kaposi's sarcomaassociated herpesvirus]	7953532 (4089, 4090) 87320849 (4091, 4092) Novel Protein sim. GBank gij4406698 gb AAD20062] - (AF131852) Unknown [Homo sapiens]	84578801 (4093, 4094) Novel Prolein sim, GBank gil4101720 (AF006466) - lymphocyte specific formin related protein [Mus musculus]		88094690 (4097, 4098) Novel Protein sim. GBank gil4589656 dbj BAA76850.1  - (AB023223) KIAA1006 protein [Homo saplens]
2   95307447 (4083, 4084)   1 (4)   1	94328076 (4085, 4086)			2047 84578801 (4093, 4094) t	2048 84606378 (4095, 4096)	88094690 (4097, 4098)

264488, 52645156, 56182575, 22278994, 35696286, 56994075, 22278998, 22278998, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 23278999, 22278999, 23278999, 23278999, 22278999, 2644259, 29331824, 265010, 264409, 264909, 265008, 6043229, 6043229, 6043229, 8108348, 265010, 87168559, 265017, 265018, 264681, 264681, 264681, 264681, 264681, 264681, 264681, 264681, 264681, 264681, 264681, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906761, 21906761, 21906761, 21906761, 21906762, 264629, 18108374, 18108370, 264628, 18108374, 18108379, 264638, 264636, 264625, 18108385, 18108338, 87168518, 264682, 264656, 26465	52646842, 52646365, 56182575, 35696286, 22278996, 22278997, 22278998, 264093, 52645060, 35696052, 29334828, 33656970, 265009, 52646317, 55111386, 52644296, 52644229, 21906769, 35695917, 265021, 60170615, 52644150, 33657109, 33697182, 27466261, 27486262, 35695763, 35696423, 35695855, 52644332	265007, 265008, 264591	. 29331825, 264682, 264686, 204691, 204093, 22279002	56182575, 28331824, 29331826, 264910, 55811957, 18108370, 55811576
synthase	UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED
Contains protein domain (PF00749) - synthase tRNA synthetases class I (E and Q)				
88177396 (4111, 4112) Novel Protein sim. GBank gil4826960[ref]NP_005042.1[pQARS - glutamine-IRNA synthetase	87877905 (4113, 4114) Novel Protein sim. GBank gij728850jspjP08640jAMYH_YEAST - GLUCOAMYLASE S1/52 PRECURSOR (GLUCAN 1,4-ALPHA- GLUCOSIDASE) (1,4-ALPHA-D-GLUCAN GLUCOHYDROLASE)	5, 4116)	7986684 (4117, 4118) Novel Protein sim. GBank gi[119714 sp P13983 EXTN_TOBAC - EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH	83050800 (4119, 4120) Novel Protein sim. GBank gij2811122 (U87318) - NaDC-2 [Xenopus laevis]
2056   88177396 (4111,	2057 87877905 (4113	2058 86276896 (4115, 4116)		2060 83050800 (4116

2061	2061   95362204 (4121, 4122)   Novel Protein sim. GBank gi 2496947 sp Q09298 YQ 141.2 KD PROTEIN EEET	Novel Protein sim. GBank gil2496947ispiQ09298jYQO9_CAEEL • HYPOTHETICAL 141.2 KD PROTEIN EEED8.9 IN CHROMOSOME II	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain		22278997, 22278999, 264259, 29331822, 29331824, 29331826, 29331827, 35696052, 29331827, 35696052, 29331828, 264908, 264906, 66712502, 29331830, 265090, 264900, 264591, 265017, 265007, 265009, 264480, 264684, 264684, 264684, 264686, 264686, 264687, 18108358, 264689, 265020, 265022, 264691, 33657109, 20281149, 18108379, 35695855, 264634, 264556, 264557, 264558, 18108382, 265526, 264557, 264558, 18108382, 264559, 83373044, 18108384, 56528488, 6043213
2062	87028440 (4123, 4124)	87028440 (4123, 4124) Novel Protein sim. GBank gil4502091  ref NP_001139.1  pANK2 - ankyrin 2, neuronal	Contains protein domain (PF00023) - struct Ank repeat		264905, 264628, 264907, 264629, 264908, 264909, 18108374, 263978, 35695855, 264512, 264635, 60431850, 264638, 264760, 264487, 264766
2063		87601272 (4125, 4126) Novel Protein sim. GBank gil4589562 dbj BAA76803.1  - (AB023176) KIAA0959 protein [Homo sapiens]	Contains protein domain (PF00617) - oncogene RasGEF domain	oncogene	22278994, 22278999, 264259, 29331827, 264906, 264909, 52644045, 264686, 21906767, 55811957, 264692, 18108365, 263972, 55811576, 18108384, 22279002, 264482, 264563, 264564, 264484
2084		95317253 (4127, 4128) Novel Protein sim. GBank gil1754515 dbj BAA13413.1 - (D87515) aminopeptidase-B [Rattus norvegicus]		hydrolase	264488, 52646365, 56994075, 35696286, 22278997, 22278998, 264259, 29331826, 2643289, 265286, 2643289, 26507, 265088, 60432229, 265097, 265008, 60432229, 265017, 265018, 265011, 287168559, 255018, 264781, 28168351, 24682, 264369, 264288, 2564429, 21906767, 21906767, 21906768, 35695917, 33657109, 18108368, 18108374, 35696423, 35695855, 52644332, 264566, 60432113, 22279000, 22279002, 264566, 264486
2065	95092238 (4128, 4130)	2065   95092238 (4129, 4130) Novel Protein sim. GBank gi 2507144 sp Q04205 TENS_CHICK - TENSIN	·	kinase	264569, 18108394, 56182181, 60432289. 29331826, 264905, 284906, 264908. 60431735, 60433356, 55811386, 85658542, 265018, 55811150, 264681, 264766, 264692, 60431528, 263974, 55810764, 35895855, 284631, 264634, 264635, 60431850, 284557, 83373044, 18108388, 22279000, 22279002
2068		85793402 (4131, 4132) Novel Protein sim. GBank gij160171 (M58295) - circumsporozoite protein [Plasmodium yoelii]	Contains protein domain (PF00096) - UNCLASSIFIED Zinc finger, C2H2 type	UNCLASSIFIED	56182375, 264259, 264906, 264764, 264288, 56182323, 264567

35696286, 22278997, 22278998, 60432049, 264259, 60432289, 60433438, 264682, 264448, 264369, 264288, 18108355, 21908765, 21906768, 265022, 33657109, 35696423, 35695855, 264558, 264404, 264563, 264488	264687	6642419, 56182515, 22278995, 22278996, 56994075, 264259, 29331822, 29331825, 29331825, 29331825, 29331825, 29331825, 29331825, 26994075, 264965, 26182435, 265009, 60433356, 87168559, 265017, 265018, 26604, 265019, 264448, 26446, 26466, 21906765, 21906767, 21906768, 21906769, 265020, 265021, 33657023, 33657109, 263976, 264555, 264557, 56182323, 83373044, 87168518, 60432113, 22278000, 222780002	35895917, 264905, 264628, 264908, 204030	18108398, 265006, 265007, 265008, 265009, 264594, 265010, 265011, 18108351, 18108354, 18108368, 18108365, 18108388, 264634, 18108385, 18108388, 18108391	22278995, 35696286, 22278997, 22278998, 22278999, 22278999, 264269, 264259, 29331822, 29331826, 35696052, 265008, 33657402, 29306754, 265011, 265019, 18108351, 264682, 264369, 21906765, 21906765, 21906766, 21906766, 21906766, 265021, 264690, 264692, 35696423, 264555, 264556, 264558, 22279000	264556	29331822, 264909, 264511, 265009, 264594, 264596, 265010, 265011, 265017, 265018, 265019, 264448, 264683, 265020, 265021, 18108370, 264632, 83373044, 264567
	UNCLASSIFIED	transcriptfactor	UNCLASSIFIED			UNCLASSIFIED	
		Contains protein domain (PF00023) - transcriptfactor Ank repeat		Contains protein domain (PF00568) • WH1 domain	Contains protein domain (PF00184) - Neurohypophysial hormones, C- terminal Domain		
		94319177 (4137, 4138) Novel Protein sim. GBank gij3152662 (AF084604) - KE03 protein [Homo sapiens]	85791380 (4139, 4140) Novel Protein sim. GBank gil5712131 gb AAD47379.1 AF12049 - (AF120499) DEM1 protein [Homo sapiens]	86946116 (4141, 4142) Novel Protein sim. GBank gij3551531 dbj BAA33016  - (AB017437) avena [Gallus gallus]		27925664 (4145, 4146) Novel Protein sim. GBank gij1504026 dbi BAA13212  - (D86976) similar to C.etegans protein (Z37093) [Homo sapiens]	94324767 (4147, 4148) Novel Protein sim. GBank gil4240317(bb)[BAA74937.1] - (AB020721) KIAA0914 protein [Homo saplens]
95303892 (4133, 4134)	84344754 (4135, 4136)	34319177 (4137, 4138)	85791380 (4139, 4140)	86946116 (4141, 4142)	91718429 (4143, 4144) ,	27925664 (4145, 4146)	
2067 9	2068		2070	2071	2072	2073	2074

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		UNCLASSIFIED	ubiquitin	UNCLASS				UNCLASS		eph			collagen				transport					UNCLASSIFIED
			Contains protein domain (PF00628) - ubiquitin PHD-finger	Contains protein domain (PF00098) - UNCLASSIFIED Zinc finger, CCHC class				Contains protein domain (PF01363) - UNCLASSIFIED	FTVE zinc imger	Contains protein domain (PF00431) - eph	CUB domain.						Contains protein domain (PF00153) - transport	Mitochondrial carrier proteins				
94314886 (4149, 4150) Novel Protein sim. GBank gi[5138930 gb AAD40362.1  - (AF093680) transcription factor IIB [Homo sapiens]			87539364 (4155, 4156) Novel Protein sim. GBank gil4220590(db) BAA74579  - (D87908) nuclear protein np95 [Mus musculus]	88095916 (4157, 4158) Novel Protein sim. GBank gil4240255 dbj BAA74906.1 -  (AB020690) KIAA0883 protein Homo saplens				94136689 (4159, 4160) Novel Protein sim. GBank gil2408021jembjCAB16219.1]	(castoz) putative vacuorar protein į schizosaccharomyces [pombe]		gl 5524734 gb AAD44360.1 AF16635 - (AF166350) ST7  protein [Homo sapiens]		87628629 (4163, 4164) Novel Protein sim. GBank gij3880558 emb CAA94234 -	(Z/0Z/1) predicted using Genefinder; similar to collagen; cDNA EST yk308e7.3 comes from this gene; cDNA EST	yk308e7.5 comes from this gene; cDNA EST yk385a8.3	comes from this gene, conv. EST yk365a6.5 comes from this gene (Caeno	94141000 (4165, 4166) Novel Protein sim. GBank gij2352427 (AF004161) -	peroxisomal Ca-dependent solute carrier [Oryctolagus	Solon			93199230 (4107, 4109) novel Protein sm. GBank gil728836 sp P39193 ALUE_HUMAN - III! ALU SUBFAMILY SP WARNING ENTRY IIII
2075 94314886 (4149, 4150) N	2076 87594118 (4151, 4152)	2077 11389877 (4153, 4154)		2079   88095916 (4157, 4158)   				2080   94136689 (4159, 4160)		2081 94847186 (4161, 4162)	•		2082 87628629 (4163, 4164)		-		2083 94141000 (4165, 4166)				2000 1000000000000000000000000000000000	93189290 (4107, 4100)

Clark EST EMBL_035149 Contestion where P71937;   Clark EST EMBL_035149 Contestion where P719344;   Clark EST EMBL_035149 Contestion while P719444;   Clark EST
Cadherin
334.1 -
cadherin
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					265022, 60170615, 33657023, 35696423,
					35695855, 264952, 18108387, 22279000
2091	95309161 (4181, 4182)	95309161 (4181, 4182) Novel Protein sim. GBank		UNCLASSIFIED	263994, 264905, 264908, 264511, 264512,
		dil4580997lqb[AAD24571,1]AF12108 - (AF121081) CAMP	-		265008, 264910, 55811386, 264288, 264768,
		inducible 2 protein [Mus musculus]			56181562, 21906765, 21906768, 21906769,
					265022, 264628, 264563, 264567
2082	88223605 (4183, 4184)			homeobox	22278997, 22278999, 66712502, 87168559,
					284683, 265021, 264486
2093	87406073 (4185, 4186)	87406073 (4185, 4186) Novel Protein sim. GBank gi[2352427 (AF004161) -	Contains protein domain (PF00153) - Iransport	transport	264905, 264906, 264907, 264908, 264510,
			Mitochondrial carrier proteins		265006, 265007, 265009, 264910, 264596,
					21906754, 87168474, 265011, 264603,
		•			265018, 265019, 264760, 264766, 264768,
				_	264769, 21906767, 21906768, 21906769.
					265021, 264690, 33657023, 264693, 264628.
					264634, 264636, 264637, 264557, 56182323,
					264564
2094	91230929 (4187, 4188)	91230929 (4187, 4188) Novel Protein sim. GBank		MHC	35696286, 265017, 265018, 265019,
		gi 4929551 gb AAD34036.1 AF15179 - (AF151799) CGI-40			18108388
		protein [Homo sapiens]			
2095		95351526 (4189, 4190) Novel Protein sim. GBank gij1363238 pir  A57284 -	Contains protein domain (PF00035) - dna_ma_bind	dna_ma_bind	35696286, 52644045, 265006, 265007,
		spermatid perinuclear RNA-binding protein Spnr - mouse	Double-stranded RNA binding motif		265008, 87168559, 18108351, 21906769,
	-				29148784, 265020, 33657023, 27486262,
					18108374, 18108388
2096	94119760 (4191, 4192)	94119760 (4191, 4192) Novel Protein sim. GBank gil3834423 (AF070689) -	Contains protein domain (PF00400) - ATPase_associated	ATPase_associated	
			WD domain, G-beta repeat		
		[Orosophila melanogaster]			60432289, 29331826, 35696052, 264107,
				•	264508, 264509, 264905, 264906, 264907,
				·	264908, 52644045, 264909, 264510, 264511,
					264512, 265008, 264910, 265009, 264592.
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			-		55812038, 21906754, 264601, 264602,
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			<i></i>	-	264638, 264639, 264563, 264483, 264565,
	,				264566, 204460, 204367

2097	95322772 (4193, 4194)	2097   95322772 (4193, 4194) Novel Protein sim. GBank   oil5174501freffNP 006051.1lpl.YF1 - zinc finger protein	Contains protein domain (PF00096) - transcriptfactor Zinc finger C2H2 tyne	transcriptfactor	65274572, 264511, 265010, 264600, 265017, 26448 26428 265011 60170815 264692
					33657109, 18108370, 264636, 264483
2098	87780340 (4195, 4196)	Novel Protein sim. GBank gil4758208 ref NP_004081.1 pDUSP - dual specificity phosphatase 3 (vaccinia virus phosphatase VH1-related)	Contains protein domain (PF00782) - phosphatase Dual specificity phosphatase, catalytic domain	phosphatase	56994075, 264259, 264288, 265020, 264563
2099	95412927 (4197, 4198)	2099   95412927 (4197, 4198) Novel Protein sim. GBank gil2695659 (AF026954) - pyruvate dehydrogenase phosphatase regulatory subunit precursor; PDPr [Bos taurus]		phosphatase	65274572, 264805, 6527444, 264691, 264638, 264555
2100	95332656 (4199, 4200 <u>)</u>	95332656 (4199, 4200) Novel Protein sim. GBank gij3881189 emb CAB16514  - (Z99281) similar to ADP-ribosylation factor; cDNA EST EMBL:C08179 comes from this gene; cDNA EST EMBL:C08337 comes from this gene; cDNA EST EMBL:C09829 comes from this gene; cDNA EST comes from this gene; cDNA EST yk4	Contains protein domain (PF00025) - nucl_recpt ADP-ribosylation factor family	nuc <u>r</u> ecpt	56182575, 22278995, 22278996, 22278997, 22278998, 60432049, 264259, 29331822, 29331824, 29331825, 29331827, 29331827, 29331827, 29331828, 2946498, 264909, 265008, 265009, 264910, 264591, 60432229, 60433356, 33657402, 264758, 21906754, 85658542, 87168474, 265017, 265018, 265019, 264681, 18108351, 264762, 264448, 264508, 21906768, 21906765, 21906767, 21906768, 21906769, 265021, 33657023, 18108374, 36596423, 264558, 83373044, 18168374, 35696423, 264568, 27279007, 2727007, 27279007, 2727007, 2727007, 2727007, 27270
2101	87762604 (4201, 4202)	87762604 (4201, 4202) Novel Protein sim. GBank gil4589468 dbj BAA76761.1 -  (AB012808) mBOCT [Mus musculus]		UNCLASSIFIED	264091, 29331824, 264105, 265007, 265010, 18108380
2102	87770461 (4203, 4204) ,	87770461 (4203, 4204) Novel Protein sim. GBank gij3874149jembjCAA97423.1j - (Z73103) predicted using Genefinder (Caenorhabditis elegans]		UNCLASSIFIED	264488, 264489, 35696286, 264259, 35696052, 264508, 264905, 264907, 264908, 264909, 264511, 264512, 264591, 264593, 60433356, 264758, 264601, 264605, 264760, 18108351, 264448, 264784, 264288, 264767, 264768, 21908769, 35695917, 18108374, 264634, 264555, 264559, 264563, 264482,
2103		95413576 (4205, 4206) Novel Protein sim. GBank gi 4240159 dbj BAA74858.1  - (AB020642) KIAA0835 protein [Homo saplens]	Contains protein domain (PF01530) - transcriptfactor Zinc finger, C2HC type	transcriptfactor	65274572, 56994075, 22278999, 264259, 29331824, 29331824, 29331825, 29596052, 29331828, 265712502, 265009, 60170831, 264595, 33109954, 85658542, 87768559, 265017, 265019, 264448, 21906765, 21906768, 265022, 33657023, 27486262, 33657349, 35695763, 60431528, 18108374, 55811576, 2648223, 18108387, 87168518, 60432113, 264564
219	85776161 (4207, 4208)			UNCLASSIFIED	264592, 264604, 22279000

264488, 56182575, 22278994, 56994075, 22278996, 22278996, 22278996, 22278998, 22278999, 22278999, 26043249, 264269, 2643249, 26331822, 29331824, 29331826, 2643269, 264500, 264510, 264511, 265007, 264512, 265009, 60170831, 60432229, 33657402, 6043336, 224595, 60433438, 264762, 18108351, 265017, 265018, 265019, 264762, 18108351, 264684, 18108354, 264288, 264688, 35695917, 265018, 265017, 265018, 264692, 365019, 264692, 366922, 60170615, 52644150, 264692, 3365703, 264693, 3695703, 264635, 60431528, 18108374, 65274791, 35695855, 264635, 6043113, 264564, 264566, 264567	265006, 265019	264906, 264639	264905, 264906, 264907, 264908, 264909, 264758, 265011, 264600, 264601, 264764,	264766, 264767, 264768, 264769, 264693, 264629, 35695855, 264632, 264634, 264636, 264638, 264638, 83373044, 264486	18108348, 264769, 18108370, 18108374, 264555, 264556, 264557, 264558	264564	264766, 35695917, 264630, 264567, 264486	264508, 264906, 264591, 264682, 22279002	265006	66714117, 29331826, 29331827, 60433438. 55812038, 265017, 265019, 264689. 21906769, 55811957, 265020, 265021. 33657109, 60170394, 264558	65274572, 264689, 264691, 264692, 60432113
UNCLASSIFIED			UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	struct	cadherin		UNCLASSIFIED
								Contains protein domain (PF00069) - struct Eukaryotic protein kinase domain	Contains protein domain (PF00028) - cadherin Cadherin domain		-
2105   94848080 (4209, 4210) Novel Prolein sim. GBank gil1707032 (U80445) - coded for by C. elegans cDNA yk13g5.3; coded for by C. elegans cDNA CENSE18F; coded for by C. elegans cDNA yk126b1.3; coded for by C. elegans cDNA yk65h8.3; coded for by C. elegans cDNA yk65h8.3; coded for by C. elegans cDNA yk65h8	83365475 (4211, 4212)	79822662 (4213, 4214)[Novel Protein sim. GBank gij3881524[emb CAA93883] - [Z70039] ZK1067.4 [Caenorhabditis elegans]	94233976 (4215, 4216) Novel Protein sim. GBank gij3176689 (AC003671) -	14 gb Z35927 from S. cerevisiae. [Arabidopsis thaliana]	80478719 (4217, 4218)	87729075 (4219, 4220) Novel Protein sim. GBank gil481043 pir  S37671 - bat2 protein - human	87818419 (4221, 4222)	87293783 (4223, 4224) Novel Protein sim. GBank gij2143639 pir   56542 - calmodulin-binding protein - rat	78941388 (4225, 4226) Novel Protein sim. GBank gil4426629 gb AAD20459  - (AF100960) protocadherin [Rattus norvegicus]	87889342 (4227, 4228) Novel Protein sim. GBank gij3327184(dbjjBAA31660j - (AB014585) KIAA0685 protein [Homo sapiens]	2115   90993785 (4229, 4230) Novel Protein sim. GBank gil4757890 ref NP_004328.1 pC8OR - chromosome 8 open reading frame 1

35696286, 22278999, 56182181, 29331824, 29331825, 29331827, 35696052, 264907, 56182435, 265008, 264591, 55812038, 55811386, 87168559, 264288, 264369, 21908769, 29148629, 33657023, 35695763, 55811576, 35696423, 18108385	29331824, 264511, 265009, 33109954, 265017, 265018, 264288, 264689, 265020, 264692, 56526486, 264482	264259, 264905, 264908, 264510, 264510, 264511, 265009, 264910, 265010, 264602, 264298, 264693, 263987, 263972, 264638, 264659	264693, 18108385	264091, 264259, 29331826, 29331828, 265017, 264604, 264288, 264685, 265020, 264691, 18108370, 55810764, 264555, 264636, 60432113	264601, 264766, 263978	18108394, 56182575, 22278997, 29331822, 29331824, 29331824, 29331825, 29331826, 29331828, 264907, 56182435, 265007, 264910, 265010, 265018, 264686, 265020, 55811576, 264555, 264637, 18108382, 89373044, 18108383, 18108384, 56528486, 264665, 264567	18108396, 264757, 265011, 18108351, 264691, 264634, 18108385	29331822, 264906, 264907, 264591, 264639, 264563	264259, 264509, 264907, 264511, 85658542. 264763, 21906765, 35695917, 264636. 264488	264488, 264489, 29331827, 35696052, 264005, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264509, 264501, 264603, 264601, 264603, 265018, 264604, 264605, 264760, 264681, 264762, 264689, 60170615, 33657023, 33657109, 55810764, 264635, 264636, 264636, 264637, 264638, 264638, 264638, 264638, 264638, 264639, 83373044, 264564, 264566
Sind	struct	glycoprotein	UNCLASSIFIED	hde	UNCLASSIFIED		struct	transport	UNCLASSIFIED	UNCLASSIFIED
				Contains protein domain (PF00017) - eph Src homology domain 2		• •	Contains protein domain (PF01363) - struct FYVE zinc finger	Contains protein domain (PF00801) - transport PKD domain		
	87788904 (4233, 4234) Novel Protein sim. GBank gi 2330021 (AF019250) - kinesin- related protein; KRP; Costal2 [Drosophila melanogaster]	87078894 (4235, 4236) Novel Protein sim. GBank gi]1079307lpiri B55573 - nuclear pore complex glycoprotein p62 - African clawed frog	86999317 (4237, 4238) Novel Protein sim. GBank gil4021407 gblAAD15748  - (AF047690) ATP-binding cassette protein M-ABC1 [Homo sapiens]	Novel Protein sim. GBank gil4885527 ref NP_005480.1 pNSP3 - novel SH2-containing Src homology domain 2 protein 3	80021375 (4241, 4242) Novel Protein sim. GBank gil4757728 ref NP_004886.1 pAGTA - angiotensin/vasopressin receptor All/AVP-like	91230931 (4243, 4244) Novel Protein slm. GBank gil4929551lgblAAD34036.1 AF15179 - (AF151799) CGI-40 protein (Homo sapiens)	86787998 (4245, 4246) Novel Protein sim. GBank gil2224551 dbj BAA20764  - (AB002303) KIAA0305 IHomo saptens	83005951 (4247, 4248) Novel Protein sim. GBank gij5689455[dbj]BAA83011.1] - (AB028982) KIAA1059 protein (Homo sapiens)	95354041 (4249, 4250) Novel Protein sim. GBank gil728331spjP39188JALU1_HUMAN - !!!! ALU SUBFAMILY J WARNING ENTRY !!!!	95084231 (4251, 4252) Novel Protein sim. GBank gil4539264 emb CAB39853.1  - (AL049495) conserved hypothetical protein (Schizosaccharomyces pombe)
88259387 (4231, 4232) [	87788904 (4233, 4234)		86999317 (4237, 4238)	2120 - 87789395 (4239, 4240) Novel Protein sim. GBank gij4885527 ref NP_005480 protein 3	80021375 (4241, 4242)	91230931 (4243, 4244)	_			
2116	2117	2118	2119	2120	2121	2122	2123	2124	2125	2126

35696286, 29331826, 35696052, 264508, 264509, 264906, 264900, 264907, 264908, 264909, 264906, 264907, 264908, 264907, 264909, 264900, 264910, 264511, 264512, 265000, 28400, 284763, 265011, 264600, 284601, 284768, 264687, 264691, 264692, 264693, 265629, 18108374, 35695855, 264632, 264639, 264567, 264639, 264639, 264567, 264639, 264639, 264567, 264639, 264563, 264564, 264567	56182575, 35696286, 56182181, 29331824, 60432289, 35696052, 264905, 264907, 66712502, 264908, 264909, 266712502, 264909, 264510, 264764, 264288, 264369, 55812038, 264686, 264634, 264637, 264637, 264634, 264634, 264633, 264637, 56182323, 264639, 18108388, 264563, 264563, 264563, 264563, 264563, 264563, 264563, 264563, 264637, 264563, 2645644, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264644, 26464, 26464, 26464, 26464, 26464, 26464, 26464, 26464, 264644	66714117, 264828, 264595, 55812038, 55811150, 55811957, 264693, 18108374, 263978, 65274791, 18108381, 83373044, 22279000	gil2649255 (AE001012) -  otein [Archaeoglobus fulgidus] gil1086886 (U41276) - Similar to Contains protein domain (PF00805) - potassium_channel 35696052, 264909, 264768, 35695917 Caenorhabdilis elegans] Pentapeptide repeats (8 copies)	Contains protein domain (PF00122) - ATPase_associated 264488, 22278999, 264259, 29331827, 29331828 35696052, 264509, 264905. 264906, 2
6000000000			UNCLASSIFIED potassium_channel	ATPase_associated
			Contains protein domain (PF00805) - Pentapeptide repeats (8 copies)	Contains protein domain (PF00122) -
81118652 (4253, 4254) Novel Protein sim. GBank gil4868435[gb]AAD31315.1 AF14323 - (AF143236) apoptosis related protein APR-2 [Homo sapiens]			95417144 (4259, 4260) Novel Protein sim. GBank gi 2649255 (AE001012) -   Conserved hypothetical protein [Archaeoglobus fulgidus] 85723065 (4261, 4262) Novel Protein sim. GBank gi 1086886 (U41276) - Similar to   potassium channel protein. [Caenorhabditis elegans]	95361096 (4263, 4264) Novel Protein sim. GBank gil5689373/dbj BAA82973.1{ - (AB028944) KIAA1021 protein [Homo sapiens]
81118652 (4253, 4254)	87414262 (4255, 4256)	95102089 (4257, 4258)		
2127		2129	2130	2132

60424179, 52646365, 52646842, 56994075, 35696286, 22278997, 22278998, 60432049, 56182181, 66714117, 60424269, 29331826, 29331828, 35696052, 264905, 264906, 264907, 66712502, 29331830, 56182435, 265006, 264512, 265008, 60431735, 60433368, 33657402, 55812038, 33109954, 21906754, 55811386, 265010, 264603, 265017, 265018, 265019, 55811150, 18108351, 264602, 264029, 56811150, 25644229, 56181562, 21906765, 21906766, 21906767, 21906767, 21906768, 21906768, 21906767, 35699917, 265021, 60170615, 3569542, 654234, 60431850, 18108337, 60432113, 22278002, 264563, 264563, 264566	56181686, 35696286, 21906754, 55811386, 265011, 265017, 18108351, 264765, 284766, 264688, 21908768, 35695917, 265020, 33657023, 264628, 35695855, 264632, 264555, 264556, 264556, 264550, 18108382, 22279002	22278999, 29331828, 35696052, 264906. 264908, 264910, 265009, 264591, 264758, 52646317, 265011, 87168559, 264601, 18108351, 264448, 264683, 264684, 254689, 18108359, 264691, 33657023, 264692, 35695763, 264629, 35695855, 264631, 264635, 264636, 264637, 56182323, 264639, 22279002, 264564	264639	264569, 264909, 33109954, 264763, 21906768, 60170394, 18108385, 264563		264905, 264910, 264591, 55812038, 55811386, 85658542, 284760, 18108351, 18108359, 55811957, 285020, 265021, 33657023, 18108364, 55811576, 83373044, 18108385, 56526486, 264482
UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	ATPase_associaled	
		Contains protein domain (PF00628) - UNCLASSIFIED PHD-finger				
95351539 (4265, 4266) Novel Protein sim. GBank gil4220489 (AC006069) - hypothetical protein [Arabidopsis thaliana]	95412697 (4267, 4268) Novel Protein sim. GBank gil3875351 jemb CAB09415  - (296047) DY3.6 [Caenorhabditis elegans]	88079813 (4269, 4270) Novel Protein sim. GBank gil5689559 db  BAA83063.1  - (AB029034) KIAA1111 protein [Homo sapiens]	84346479 (4271, 4272) Novel Protein sim. GBank gij2662167[dbj BAA23715  - (AB007903) KIAA0443 [Homo saplens]	87637716 (4273, 4274) Novel Protein sim. GBank gil4884110 emb CAB43262.1  - (AL050090) hypothetical protein [Homo sapiens]	87395446 (4275, 4276) Novel Protein sim. GBank gil5174779lgb[AAD40696.1] - (U87804) 50 kDa protein [Caulobacter crescentus]	94843882 (4277, 4278) Novel Protein sim. GBank gij3850821jemb CAA77135  - (Y18350) Uz snRNP auxiliary factor, large subunit [Nicotiana plumbaginifolia]
2133 95351539 (4265, 4266) N	2134 95412697 (4267, 4268)	2135 88079813 (4269, 4270)	2136 84346479 (4271, 4272)	2137 87637716 (4273, 4274)	2138 87395446 (4275, 4276)	2139   94843882 (4277, 4278)

264489, 264259, 29331824, 264104, 264109, 264509, 265006, 264759, 265018, 26448, 264288, 21906768, 55811957, 265021, 33657023, 27488265, 35896423, 264636, 264559, 264556	265020, 264693	263978	22278997, 29331827; 264907, 265020, 60432113	65274572, 264259, 28331824, 28331827, 264906, 264908, 264591, 265011, 87168559, 264600 264010, 264010, 264010	20400, 203018, 204208, 204106, 21900103, 21906767, 55811576, 35696423, 65274791, 22279002	263978	264909, 60433356, 264686		264907, 264768, 264769, 18108385	264593	29331824, 29331826, 35696052, 264758,	87168474, 265018, 52644150, 33657109	22278998, 29331822, 29331824, 29331828, 284764, 264769, 21906766, 264486	22278996, 56994075, 22278999, 60432049,	264259, 29331822, 29331824, 29331828, 35696052, 29331828, 264508, 264511,	60433356, 264758, 264598, 33109954, 60174639, 265010, 265011, 87168559,	265017, 265018, 265019, 264448, 264288,	265020, 60170615, 33657109, 33657182.	33657349, 18108370, 264635, 264557, 60170394, 18108385, 87168518, 22279000	18108397, 18108398, 265007, 264591, 265011, 18108351, 18108368, 18108374, 18108388
UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED		ONCLASSIFIED	UNCLASSIFIED	alycoprotein		UNCLASSIFIED			·				kinase
											Contains protein domain (PF00059) - alycoprotein	Lectin C-type domain								
2140   87645655 (4279, 4280)   Novel Protein sim. GBank gil4417293 gb AAD20418  -   (AC007019) unknown protein (Arabidopsis thallana)			94140051 (4285, 4286) Novel Protein sim. GBank gil2135766 pirl S53362 - mucin   5AC (clone JER47) - human (fragment)	94320114 (4287, 4288) Novel Protein sim. GBank gil2078483 (U43200) - antifreeze glycopeptide AFGP polyprotein precursor [Boreogadus	Saldaj		87010515 (4291, 4292) Novel Protein sim. GBank gij1255871 (U53341) - short region of weak similarity to bovine membrane receptor p53	(PIR:S28503) [Caenorhabditis elegans]	80432911 (4293, 4294) Novel Protein sim. GBank gil3080398 emb CAA18718.1  - (AL022603) putative protein (Arabidopsis thaliana)		Novel Protein sim. GBank	gil 19863 spip 20693 jFCEZ_MOUSE - LOW AFFINITY IMMUNOGLOBULIN EPSILON FC RECEPTOR (LYMPHOCYTE IGE RECEPTOR) (FC-EPSILON-RII) (CD23)	94140059 (4299, 4300) Novel Protein sim. GBank gij5420387[embjCAB46679.1] - (AJ243459) proteophosohogiycan [Leishmanla malor]	95353241 (4301, 4302) Novel Protein sim. GBank gil5689407idbilBAA82987.11 -	(AB028958) KIAA1035 protein [Homo sapiens]					2152 79321640 (4303, 4304) Novel Protein sim. GBank gil3452473 (AF084205) - serine/Ihreonine protein kinase TAO1 [Raltus novegicus]
87645655 (4279, 4280)			94140051 (4285, 4286)					_					-							79321640 (4303, 4304)
2140	2141	2142	2143	2144		2145	2148		2147	2148	2149		2150	2151						2152

264488, 263994, 52646842, 22278996, 22278998, 22478999, 264259, 29331822, 35696052, 264509, 264509, 264905, 264906, 264907, 264510, 264510, 264511, 264512, 264758, 87168474, 87168559, 265017, 265019, 264769, 524768, 264768, 264769, 264769, 264769, 264687, 264769, 264687, 264691, 264632, 264635, 264639, 18108385, 264683, 264639, 18108385, 264483, 264635, 264636, 264639, 18108385, 264483, 264584, 264636, 264646, 264646, 264646, 264646, 264646, 264646, 264646, 264646, 264646, 264646, 264646,			265007, 284684		۵				IED (55274572, 264508, 264905, 264906, 264907, 264908, 52644045, 264909, 265007, 264910, 264591, 264592, 264593, 55812038, 264596,	264758, 265011, 264600, 264762, 264763.	264768, 264769, 264689, 265020, 264691,	264628, 264629, 263978, 264632, 264634.	264557, 264639, 264639, 18108385, 264563, 264566, 264567	FED 56182575, 22278998, 264093, 264683, 33657023, 65274620, 60432113		264603, 264637, 264565
glycoprotein	UNCLASSIFIED	UNCLASSIFIED			UNCLASSIFIED	dna_rna_bin		UNCLASSIFIED	UNCLASSIFIED					UNCLASSIFIED		- collagen
Contains protein domain (PF00560) - glycoprotein Leucine Rich Repeat						Contains protein domain (PF00651) - dna_rna_bind BTB/PO2 domain									82	Contains protein domain (PF01006) - collagen Hepatitis C virus non-structural
D	87408034 (4307, 4308) Novel Protein sim. GBank gi 225150 pr  1209265U - chorion protein B11 [Bombyx mori]		84295205 (4311, 4312) Novel Protein sim. GBank gij3970966 (AC004974) - spa-1- like: similar to AF026504 (PID:02555183) (Homo sapiens)		8644218 (4315, 4316) Novel Protein sim. GBank gi]1076211 pir  S50755 - https://doi.org/10/10/10/10/10/10/10/10/10/10/10/10/10/	80083729 (4317, 4318) Novel Protein slm. GBank gil4650844 dbj BAA77027.11- [AB026190] Kelch motif containing protein [Homo sapiens]	16283674 (4319, 4320) Novel Protein sim. GBank gi 2879925 dbj BAA24826  - (AB007897) KIAA0437 [Homo saplens]	-	94319526 (4323, 4324) Novel Protein sim. GBank gi[1504006 dbj]BAA13202  - (D86966) similarto human ZFY protein. [Homo sapiens]	·				95417158 (4325, 4326) Novel Protein sim. GBank gij3876537[emb[CAA98270]	(27.95/4) CONT. EST VA2916.5 Comes from this gene [Caenorhabditis element]	
18313371 (4305, 4306)	87408034 (4307, 4308)	87424072 (4309, 4310)	84295205 (4311, 4312)	87316344 (4313, 4314)	86444218 (4315, 4316)	80083729 (4317, 4318)	16283674 (4319, 4320)	87739131 (4321, 4322)	94319526 (4323, 4324)					_		80569456 (4327, 4328)
1153	2154	2155	2156		2158	2159	2160	2161	2162					2163		2164

2165	94329169 (4329, 4330)	2165   94329169 (4329, 4330) Novel Protein sim. GBank gi 1086794 (U41107) - No		UNCLASSIFIED	56994075, 22278996, 22278997, 22278999,   264250, 20111822, 20111824, 20111825
		deminion line tound   Caenomagains elegans			29331826, 29331827, 29331828, 264906,
					29331830, 56182435, 265009, 21906754,
	<u></u>			•	33657084, 265011, 265019, 264448, 264288,
		-			264369, 21906765, 21906768, 21906769,
					265020, 265021, 264691, 264692, 33657023,
	-				65274620, 35695855, 264556, 60170394,
					83373044, 60432113, 22279002, 264567
2168	87618934 (4331, 4332)	87618934 (4331, 4332) Novel Protein sim. GBank gil2706522lemb CAA75816  -		ubiquitin	52645156, 22278994, 22278998, 66714117.
_		(Y15895) ubiquitin activating enzyme [Drosophila			29331828, 52644045, 265018, 265019,
		melanogasteri			264369, 21906765, 21906767, 21906768,
					21906769, 265021, 265022, 264693,
					27486262, 35695763, 18108376, 56526486,
					8/168518, 26455/
2167	87716864 (4333, 4334,	87716864 (4333, 4334) Novel Protein sim. GBank gij2224713/dbjjBAA20840 -		UNCLASSIFIED	301023/3, 33030200, 23331024, 23331020,
		(ABUUZ384) KIAAU386 [Homo sapiens]			28140430, 30102433, 203000, 203003, 364603 364603 33657403 33100054
					264554, 204333, 33031404, 33133333, 364369
_					24006764 3400676F 3400876B 30148637
					21906/04, 21906/03, 21906/06, 29140627,
					21906/69, 52644150, 53657103, 53696423,
					18108381, 18108384, 18108385, 60432113,
					264567
2168	86999334 (4335, 4336	86999334 (4335, 4336) Novel Protein sim. GBank gil4321407 gb AAD 15748  -	Contains protein domain (PF00564) - transport	transport	66714117, 29331827, 264907, 264511,
		(AF047690) ATP-binding cassette protein M-ABC1 [Homo	ABC transporter transmembrane		264591, 265018, 264764, 264683, 264766.
		sapiens]	region.		284768, 264568
2169					264629, 264555, 264559
2170		94141033 (4339, 4340) Novel Protein sim. GBank		UNCLASSIFIED	65274572, 56182575, 22278997, 22278998,
		qi 5106521 ab AAD39741.1 AF10536 - (AF105365) K-CI			264259, 29331825, 264509, 264906,
					56182435, 60433438, 55812038, 264596,
					55811386, 265019, 264762, 264763, 264448,
					264764, 264684, 264288, 264766, 264685.
					56181562, 264689, 55811957, 265020.
	-				264535, 264691, 33657109, 60431528.
					18108374, 35696423, 55811576, 65274791,
					284634, 264639, 264558, 87168518,
					60432113, 264564
2171	80194050 (4341, 4342)	(2)		UNCLASSIFIED	264369, 265020, 264558
2172	_	(‡		UNCLASSIFIED	264259, 264558
2173		87036740 (4345, 4346) Novel Protein slm. GBank gij4309681jgbJAAD15478j -		UNCLASSIFIED	264369
	_	(AC006930) R33423_1 [Homo sapiens]			
2174	_	95003288 (4347, 4348) Novel Protein sim. GBank			264906, 35695855, 264555, 264557
		gij2493778jspjQ09456jYQ35_CAEEL - PUTATIVE			
		COLLEGE COLLAGER COSCS.3			

	18108392, 222788997, 222788999, 2640993, 33657402, 265019, 264448, 264766, 264689, 21906767, 21906768, 21906769, 265021, 33657023, 18108370, 18108374, 60432113, 22279002		UNCLASSIFIED 60424269, 264760, 264628, 264632
oncogen	kinase	- (ebh	UNCLA
Contains protein domain (PF00805) Penlapeptide repeats (8 copies)		s Contains protein domain (PF00012 Hsp70 protein	
	Novel Protein sim. GBank gi[5454072 ref NP_006416.1 pSLU7 - step II splicing factor SLU7		
88223392 (4351, 4352)	2177   94128942 (4353, 4354) ,	2178 87601557 (4355, 4356)	2470 87246276 (4267 4268)
	Contains protein domain (PF00805) - oncogene	Contains protein domain (PF00805) - oncogene 7_HUMAN - III ALU SUBFAMILY Pentapeptide repeats (8 copies) 6.1jpSLU7 - step II splicing factor	Contains protein domain (PF00805) - oncogene 7_HUMAN - III ALU SUBFAMILY Pentapeptide repeats (8 copies) 6.1pSLU7 - step II splicing factor gil473407 (U08215) - NST-1 [Mus Contains protein domain (PF00012) - eph Hsp70 protein

52644507, 22278994, 35696286, 22278997, 22278999, 264259, 52645080, 29331822, 29331824, 29331824, 29331824, 29331824, 29331824, 29331824, 29331824, 29331826, 29331826, 29331826, 29331826, 29331830, 264909, 264510, 265006, 265007, 33657402, 55612038, 21906754, 87168474, 87168559, 265017, 265019, 265019, 264686, 21906765, 21906766, 21906769, 265020, 265021, 265022, 52644150, 33657023, 33657109, 27486265, 33657349, 18108374, 35696423, 35656486, 87168518, 60432113, 22279000, 264482, 264568, 264567, 264686, 264568, 264567, 264486	29331827, 264369, 16108376, 204364	22278996, 22278997, 22278999, 264259, 29331822, 56182435, 264112, 264764, 264288, 21906767, 21906768, 21906769, 33657109, 18108376, 60170394, 22279000, 22279002	264760	29331822, 29331825, 29331826, 56182435, 265011, 264685, 264686, 21906768, 18108370, 264629, 264631, 264639, 264557	29331824, 264907, 66172502, 264757, 265019, 264288, 264692, 56526486	ATPase_associated   264259, 29331822, 29331824, 29331826, 56182436, 264592, 55812038, 264760, 264766, 55811957, 33657023, 55811576, 56182323, 264563	35696052, 264905, 264906, 264907, 264908, 264510, 264511, 265008, 264910, 264758, 265019, 264762, 264681, 264766, 264769, 35695917, 264692, 35695423, 264631, 264635, 264637, 18108388, 264566, 264486	18108398, 56994075, 264259, 29331824, 29331825, 66714117, 29331827, 264906, 29331830, 265018, 265020, 265021, 56182323, 264559, 22279000, 22279002	22278996, 264259, 29331826, 21906754, 264369, 264288, 263967
- kinase	kinase	UNCLASSIFIED		tnf		ATPase_associate	UNCLASSIFIED	histone	- (0
Contains protein domain (PF00400) - kinase WD domain, G-beta repeat				`		·			Contains protein domain (PF01596) - O-methyltransferase
HB_DICDI - MYOSIN HEAVY B)	85764930 (4361, 4362) Novel Protein sim. GBank gi 3024689 sp Q15542 T2D4_HUMAN - TRANSCRIPTION INITIATION FACTOR TFIID 100 KD SUBUNIT (TAFII-100)  (TAFII100)	Novel Protein sim. GBank giļ5420387 emb CAB46679.1  - (AJ243459) proteophosphoglycan [Leishmanla major]	85460649 (4385, 4366) Novel Protein sim. GBank gij3873406jgbjAAC77482.1  - (U17129) unknown IRhodococcus erythropolisi	87760690 (4367, 4368) Novel Protein sim. GBank gij3114713 (AF061346) - Edp1 protein [Mus musculus]	87826463 (4369, 4370) Novel Protein sim. GBank gl 5106956 gb AAD39906.1 AF11361 - (AF113615) FH1/FH2 domain-containing protein FHOS [Homo sapiens]	87739227 (4371, 4372) Novel Protein slm. GBank gilz864625jemb CAA16972  - (AL021811) putative protein [Arabidopsis thaliana]		87771708 (4375, 4376) Novel Protein sim. GBank gils 10781619413 contains gils 107816194040129.1 pt 14941 - (AF149413) contains similarity to histone deacetylases; Pfam PF00850. Score=13.3, E=5e-10, N=1 [Arabidopsis thaliana]	85693573 (4377, 4378) Novel Protein sim. GBank gij3452357 (AF075724) - unknown [Legionella pneumophila]
95351397 (4359, 4360)	85764930 (4361, 4362)						87388173 (4373, 4374)		
2180	2181	2182	2183	2184	2185	2186	2187	2188	2189

06	87639197 (4379, 4380)			nucleaseinhib	22278996, 22278999, 29331822, 29331824, 29331826, 265008, 264910, 60170831, 55812038, 52844298, 265010, 265018, 284685, 264688, 56181562, 21906769, 35695917, 265022, 60170394, 22279000
	95198928 (4381, 4382)	95198928 (4381, 4382) Novel Protein sim. GBank gij5327002jemb CAB46272.1  - (Y18503) XAP-5-like protein [Homo sapiens]		·	29331825, 29331826, 29331830, 264510, 264511, 264910, 264593, 264594, 264556, 264559
	11126316 (4383, 4384)	11126316 (4383, 4384) Novel Protein sim. GBank  gi 462600 sp P34400 Mi10_CAEEL - MIG-10 PROTEIN	Contains protein domain (PF00169) - PH domain		264558
	94140073 (4385, 4386)	Novel Protein sim. GBank gil\$420389[emb CAB46680.1  - (AJ243460) proteophosphoglycan [Leishmania major]		UNCLASSIFIED	56181686, 29331825, 29331827, 264508, 264909, 264909, 265008, 264592, 60432229, 264288, 264684, 264766, 35695917, 33657023, 60431602, 60431528, 55810764, 55811576, 65274781, 35695855, 60431850, 56182323, 60432113
	21418714 (4387, 4388)	21418714 (4387, 4388) Novel Protein sim. GBank gil2773341 (AF040954) - putative protein phosphatase 1 nuclear targeting subunit [Rattus norvegicus]			264592
	88083023 (4389, 4390)	88083023 (4389, 4390) Novel Protein sim. GBank gi[2832763 emb]CAA15685.1  - (AL009191) /prediction=(method;; /prediction=(method;; //match=(desc;; /match=(desc;; /motif=(desc; [Orosophila melanogaster]		UNCLASSIFIED	22278996, 22278999, 35696052, 265006, 21806754, 265017, 35695917, 265021, 265022, 35695855
	95091631 (4391, 4392)	Novel Prolein sim. GBank gi[5262487 emb[CAB45699.1] - (AL080076) hypothetical protein [Homo sapiens]		collagen	56182575, 35696286, 22278997, 22278999, 264259, 29331822, 66714117, 60432289, 29331827, 35696052, 29331828, 264508, 25244045, 56182435, 264510, 265007, 265008, 265009, 60433439, 55112038, 265010, 265011, 26448, 264288, 264686, 264687, 5264229, 21906765, 21906765, 21906767, 35695917, 265022, 264691, 356954023, 264693, 18108370, 18108376, 35695423, 55811576, 65274791, 35695855, 265638, 56182323, 18108388, 265638, 56182323, 18108388, 265638, 56182323, 18108388, 265638, 56182323, 18108388, 265638, 56182323, 18108388, 265638, 56182323, 18108388, 265638, 56182323, 18108388, 265638, 56182323, 18108388, 265638, 56182323, 18108388, 265638, 56182323, 18108388, 265638, 56182323, 18108388, 265638, 56182323, 18108388, 265638, 56182323, 561823825, 56182223, 56182223, 56182223, 56182223, 56182223, 56182223, 56182223, 561822223, 561822223, 561822223, 5618222222222222222222222222222222222222
	95073813 (4393, 4394)	95073613 (4393, 4394) Novel Protein sim. GBank gil4929567[gb]AAD34044.1 AF15180 - (AF151807) CG1-49 protein [Homo sapiens]			264768, 264769, 21906765, 21906766, 21906767, 29148627, 55811957, 35696286, 265020, 22278998, 265021, 264259, 33657023, 264693, 29331824, 35696052, 29331828, 18108370, 35695855, 264113, 265008, 264910, 60432229, 56182323, 33657402, 264758, 83333044, 21906754, 265018, 265019, 22279002, 264482, 264448, 24465, 264288, 244369
2198	88060914 (4395, 4396)	88050914 (4395, 4396) Novel Protein sim. GBank gi 3548787 (AC005622) - R30953_1 (Homo sapiens)		UNCLASSIFIED	

2199	88054355 (4397, 4398)	2199   88054355 (4397, 4398)   Novel Protein sim. GBank gi[2739372 (AC002505) -			264105, 264110, 264112, 264688, 55811957,
		nypomencal protein (Arabidopsis manaria)			55526486
2200	87405385 (4399, 4400)	87405385 (4399, 4400) Novel Protein sim. GBank gij3043634[dbjjBAA25481] - (AB011127) KIAA0555 protein [Homo sapiens]		struct	29331824, 264763, 264768
2201	94316872 (4401, 4402)	94316872 (4401, 4402) Novel Protein sim. GBank. gij3913470[sp]057314[DHBX_ANAPL - PUTATIVE STEROID DEHYDROGENASE SPM2	Contains protein domain (PF00106) - dehydrogenase short chain dehydrogenase		29331824, 35696052, 264905, 264907, 33657402, 55811386, 265017, 265018, 265019, 265019, 265022, 33657023, 33657109, 27486261, 18108370, 35696423, 3569585, 264555, 264556, 83373044, 87168518,
2202	91672385 (4403, 4404)	91672385 (4403, 4404) Novel Protein sim. GBank gi 5262655 emb CAB45767.1  - (AL080186) hypothetical protein [Homo sapiens]		UNCLASSIFIED	264489, 264259, 29331824, 60432289, 265052, 265017, 265062, 26439, 264592, 265017, 265018, 265018, 265018, 265018, 265018, 265018, 265018, 264288, 264766, 21906765, 21906766, 264690, 264691, 264692, 33557109, 264659, 264636, 264555, 264639, 264558, 264559, 83373044, 18108385, 264404, 22279002, 264482
		87761832 (4405, 4406) Novel Protein sim. GBank gij1172845 sp P46629 RB25_RABIT - RAS-RELATED PROTEIN RAB-25	Contains protein domain (PF00071) - glycoprotein Ras family	glycoprotein	52646365, 56994075, 264259, 29331822, 29331826, 29331822, 29331827, 29331828, 264910, 265010, 265010, 265019, 264605, 264288, 21906769, 35695917, 33657023, 264638, 22279000, 264566, 264567
2204		88088671 (4407, 4408) Novel Protein sim. GBank gij121036 sp P29348 GBT3. RAT   Contains protein domair GUANINE NUCLEOTIDE-BINDING PROTEIN G(T), ALPHA-G-protein alpha subunit 3 SUBUNIT (GUSTDUCIN ALPHA-3 CHAIN)	Contains protein domain (PF00503) - UNCLASSIFIED G-protein alpha subunit	UNCLASSIFIED	
2205		94147589 (4409, 4410) Novel Protein sim. GBank gil4589480 db  BAA76768.1  - (AB023141) KIAA0924 protein [Homo saplens]	Contains protein domain (PF00096) - dna_rna_bind Zinc finger, C2H2 type	dna_rna_bind	18108394, 18108397, 56182575, 60432049, 264259, 29331822, 29331824, 29331825, 29331826, 29331826, 29331826, 29331827, 264806, 265007, 265008, 264680, 264308, 264369, 264696, 264691, 264691, 264693, 18108388, 5811576, 264631, 264693, 18108381, 18108384, 60432113, 22279002, 264563, 264566
2206				UNCLASSIFIED	264591
2207		87787970 (4413, 4414) Novel Protein sim. GBank gil4557753 ref NP_000372.1 pMID1 - midline 1 protein	Contains protein domain (PF00622) - SPRY domain		29331822, 56182181, 29331827, 35696052, 52644045, 265006, 265019, 56181562, 55811957, 265021, 33657023, 35695763, 35695855, 60170394, 60432113, 264568
2208	_				264908, 265019, 18108351, 21906769
5209		87800420 (4417, 4418) Novel Protein sim. GBank gij3986746 (AF105228) - tuftelin [[Bos taurus]		struct	264112, 285009, 264691, 18108385, 18108374, 264634, 20281166

210	57152407 (4419, 4420)	2210   57152407 (4419, 4420)   Novel Protein sim. GBank		kinase	284803
		gii/zb837(spip39194(ALU7_HUMAN - !!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!			
2211	87341720 (4421, 4422)	2211 (87341720 (4421, 4422) Novel Protein sim. GBank		oncogene	264685, 264686, 18108365, 22279002,
		gif28833[spjP39194JALU7_HUMAN - IIII ALU SUBFAMILY SO WARNING FNTRY IIII	٠		264482
2212	91223924 (4423, 4424)	91223924 (4423, 4424) Novel Protein sim. GBank gij3776027 emb CAA09214  -	Contains protein domain (PF00270) - helicase	helicase	22278995, 22278997, 22278999, 264092,
		(AJ010475) RNA helicase [Arabidopsis thaliana]	DEAD/DEAH box helicase		264094, 29331822, 66714117, 29331826,
					29331828, 264907, 52644045, 265009,
					60170831, 21906754, 87168559, 265017,
		•			265019, 18108351, 264683, 18108354,
					264369, 264766, 264687, 52644229,
					21906765, 21906766, 21906767, 21906768,
			-		265021, 33657109, 18108370, 18108374,
					264638, 56182323, 18108384, 18108387,
					87168518, 264565
2213	91219309 (4425, 4426)	2213  91219309 (4425, 4426) Novel Protein sim. GBank gil5420387 lemb (CAB46679.11 -			56182575, 22278996, 22278997, 35696052,
		(AJ243459) proteophosphoglycan [Leishmania major]	•		264905, 66712502, 264908, 264828,
				•	56182435, 264112, 265008, 60431735,
					60433438, 21906754, 265010, 265011,
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					264558, 83373044, 22279002, 264482,
					264483

264488, 5264507, 16108394, 56182575, 22278994, 22278994, 22278996, 2694076, 22278997, 22278999, 22696286, 56994076, 22278997, 22278999, 22278999, 264490, 60432049, 264259, 52645080, 28331822, 29347620, 29331824, 2645080, 26331826, 29331827, 3569652, 29331828, 20281100, 264509, 264907, 66772502, 264908, 26331826, 20281100, 264509, 264511, 265007, 265008, 265009, 60170831, 264519, 265007, 265008, 2650010, 265011, 87166559, 264601, 265010, 265011, 265011, 265013, 26448, 264682, 264682, 264687, 265012, 265021, 265022, 264632, 21906766, 21906766, 21906766, 21906766, 21906766, 21906767, 21906768, 21906769, 33657023, 264692, 18108364, 3365703, 264692, 27486264, 27486265, 35695855, 264634, 264636, 52644332, 264638, 264634, 18108384, 18108376, 264636, 52644332, 264638, 264634, 18108381, 265528, 18108387, 18108381, 264538, 264538, 18108381, 18108384, 18108381, 18108384, 18108386, 18108387, 18108388, 18108387, 18108388, 18108387, 18108388, 18108387, 18108388, 181	222/19UUZ, 26482, 264564, 264566, 264966, 264909, 265008, 264555, 264558, 87168518	264693	284288, 33657109, 264556	35696423, 264563	264682, 264683, 264688, 264689, 264693, 18108370, 18108376
dehydrogenase		UNCLASSIFIED	glycoprotein		
Contains protein domain (PF00725) - dehydrogenase 3-hydroxyacyl-CoA dehydrogenase			Contains protein domain (PF00560) - glycoprotein Leucine Rich Repeat	Contains protein domain (PF00069) - Eukaryotic protein kinase domain	Contains protein domain (PF01963) - TraB family
95361453 (4427, 4426) Novel Protein sim. GBank gil4564325 reflNP_000173.1 pHADH - hydroxyacyl- Coenzyme A dehydrogenase/3-ketoacyl-Coenzyme A thiolase/enoyt-Coenzyme A hydratase (trifunctional protein), alpha su			80589404 (4433, 4434) Novel Protein sim. GBank gi 5031707 ref NP_005503.1 pGARP - glycoprotein A repetitions predominant	85518254 (4435, 4436) Novel Protein sim. GBank gij3878636jembjCAA88953j - (Z49128) similar to cAMP-dependant protein kinase; cDNA EST EMBL.T00719 comes from this gene; cDNA EST yk46548.3 comes from this gene; cDNA EST yk46548.5 comes from this gene; cDNA EST yk46548.5 tomes from this gene; cDNA EST yk492f4.3 comes from this gene; cDNA EST yk492f4.3 comes from this gene; cDNA EST yk492f4.3 comes from	

2220	95354165 (4439, 4440)	2220   95354165 (4439, 4440)   Novel Protein sim. GBank   gi 4507261 ref NP_003145.1 pSTAT - statherin			264488, 18108394, 18108395, 35696286, 264259, 264097, 60432289, 264509, 264905, 264905, 264905, 264907, 265906, 264907, 265907, 26331830, 264908, 264909, 265009, 264510, 265009, 264511, 265007, 264512, 264910, 265009, 264593, 264594, 60433356, 264760, 26401, 264603, 265019, 264605, 264760, 264764, 264369, 264768, 264691, 18108358, 264628, 264692, 1810836, 264628, 264629, 18108374, 263378, 264634, 264638, 18108385, 264634, 264638, 264637, 264638, 18108385, 264634, 264638, 18108385, 264634, 264638, 18108385, 264634, 264638, 18108385, 264634, 264638, 18108385, 264634, 264638, 18108385, 264634, 264638, 18108385, 264634, 264638, 264657, 264638, 18108385, 264634, 264644, 264644, 264644, 264644, 264644, 264644, 264644, 264644, 264644, 2646
2221	88060927 (4441, 4442)	88060927 (4441, 4442) Novel Protein sim. GBank gij3549154 (AC005625) - R27328_1 [Homo sapiens]			
2222	2222 84425892 (4443, 4444)			UNCLASSIFIED	264908_265020_35695855
2223	95091649 (4445, 4446)			UNCLASSIFIED	265010, 264685, 264690, 264693, 264638, 263974, 263976, 55611576, 264555, 264638, 8337304, 264883
2224		87388515 (4447, 4448) Novel Protein sim. GBank gij3876005 emb CAA84799] - (Z35719) cDNA EST EMBL:D67419 comes from this gene; cDNA EST EMBL:C13853 comes from this gene; cDNA EST EMBL:C11578 comes from this gene; cDNA EST yk234a7.3 comes from this gene; cDNA EST comes from this gene; cDNA EST	Contains protein domain (PF01958) - UNCLASSIFIED Domain of unknown function	UNCLASSIFIED	264259, 264509, 56182435, 265006, 265008, 265009, 265009, 284737, 21906754, 18108351, 264693, 18108374, 18108385
2225	85749484 (4449, 4450)	85749484 (4449, 4450) Novel Protein slm. GBank gi]1255847 (U53338) - C05E11.1 gene product [Caenorhabdilis elegans]		transport	22278994, 22278995, 22278999, 52644045, 264600, 265019, 21906765, 21906769
9777	86978953 (4451, 4452)	86978953 (4451, 4452) Novel Protein sim. GBank gil4826524 emb CAB42852.1  - (AL049848) hypothetical protein [Homo sapiens]			264259, 29331822, 29331824, 29331825, 29331827, 264508, 264906, 265007, 264691, 264634, 264486
2221				UNCLASSIFIED	22278999, 265006, 265008, 18108354, 29148629, 29148784, 27486261, 18108374, 264637, 18108384
8222		91227337 (4455, 4456) Novel Protein sim. GBank gi 606976 (U16800) - ribonucleoprotein [Xenopus laevis]	Contains protein domain (PF00076) - dna_ma_bind RNA recognilion motif. (a.k.a. RRM, RBD, or RNP domain)	dna_ma_bind	264091, 264092, 264094, 29331822, 29331825, 68714117, 264693, 263972, 264639, 83373044, 264563
6777		88050931 (4457, 4458) Novel Protein sim. GBank gij3549155 (AC005625) - R27328_2 [Homo sapiens]		UNCLASSIFIED	

	IED 264563	dor 18108394, 56182575, 22278995, 35696286, 22278997, 22278999, 264259, 29331827, 35696052, 264907, 56182435, 265006.	265007, 265008, 264910, 264758, 55812038, 264603, 265018, 265019, 18108351, 264682, 264764, 264683, 264369, 264288, 264686, 264687, 264689, 21906765, 21906767, 21906767, 2148697, 2148697, 2148697, 21486769, 364687	264690, 52644150, 264691, 33657023, 264693, 18108370, 18108374, 55811576,	250930530, 2545053, 10100305, 204204 264906, 33657405, 265018, 264288, 264686,	26182435, 264369, 264688, 21906765, 265020, 264693, 264568, 56526486	IED 22278997, 264563	264569, 264687, 264769, 265022, 264259, 60432049, 264691, 29331826, 60432289, 20281149, 264906, 264907, 264511, 265008, 265009, 264634, 264635, 264636, 264556, 264557, 264558, 60433356, 264595, 264559, 60433438, 60432113, 264761,	264762, 264783, 264764 264488, 264769, 21906765, 21906766, 21906767, 21906768, 21906769, 22278995, 22278996, 22278997, 22278998, 265020, 265021, 22278999, 264259, 29331824, 29331826, 29331827, 36695763, 19108376, 35695855, 265007, 60432229, 33657402, 60433356, 60433438, 83373044, 18108385, 21906754, 18108387, 60432113, 22279000, 265019, 22279002, 264482, 18108351,
UNCLASSIFIED	UNCLASSIFIED	transcriptfactor				kinase	00) - UNCLASSII	- eph	)) - kinase
				-			Contains protein domain (PF00400) - UNCLASSIFIED WD domain, G-beta repeat	Contains protein domain (PF00011) - leph Hsp20/alpha crystallin family	Contains protein domain (PF00400) - kinase WD domain, G-beta repeat
2230   95342915 (4459, 4460) Novel Protein sim. GBank gil226154ipri[i1412350A - DNA polymerase [Human adenovins type 2]	88060937 (4461, 4462) Novel Protein sim. GBank gi[3549154 (AC005625) - R27328_1 [Homo sapiens]	Novel Protein sim, GBank gij5281316jgbfAAD41476.1 AF13312 - (AF133124) transcription factor IIIC63 (Homo sapiens)			87755292 (4465, 4466) Novel Protein slm. GBank gil4249733jgb AAD13780  -   (AF109377) IdIBp [Mus musculus]	87771817 (4467, 4468) Novel Protein sim. GBank gif1706559jspIP54352 EAS_DROME - ETHANOLAMINE KINASE (EASILY SHOCKED PROTEIN)	91012316 (4469, 4470) Novel Protein sim. GBank gil4972734 gb AAD34762.1 - (AF132174) unknown [Drosophila melanogaster]	protein - human	91012318 (4473, 4474) Novel Protein sim. GBank gil4972734lgb AAD34762.1  . (AF132174) unknown [Drosophila melanogaster
230 95342915 (4459, 4460		2232   87762581 (4463, 4464					2235 91012316 (4469, 4470) 2236 88003131 (4471, 4472)		2237   91012318 (4473, 44 <u>74)</u>

Contains potein drawn (PT00266)   244502, 284502, 284519, 7846544, 784564   First   Contains potein drawn (PT00179)   City   Contains potein drawn (PT00179)   City   Ci
[2982311 (AF051240) - Ig enzyme E2 [Picea mariana] a GAMMA SUBUNIT (EIF-2B TOR)  [2291143 (AF016417) - Similar Caenorhabditis elegans] 470340 (U00043) - similar to aenorhabditis elegans]
[2982311 (AF051240) - Ig enzyme E2 [Picea mariana] a GAMMA SUBUNIT (EIF-2B TOR)  [2291143 (AF016417) - Similar Caenorhabditis elegans] 470340 (U00043) - similar to aenorhabditis elegans]

246	94848710 (4481, 4492)	2246 94848710 (4481, 4482) Novel Protein sim. GBank gil4996096jdbjjBAA78326.11 -	Contains protein domain (PF00153) - Iransport		65274572, 22278995, 35696286, 22278996,
					264106, 264905, 264907, 265006, 265007, 265008, 60433438, 33109954, 87168559,
					265018, 265019, 264288, 21906765, 21906767, 21906767, 21906768, 21906769, 55811957,
					35695917, 265020, 265022, 27486264,
					18108370, 18108374, 65274791, 35695855. 60432113
2247	87862542 (4493, 4494)	87862542 (4493, 4494) Novel Protein sim. GBank gil854065[emb[CAA58337] -		UNCLASSIFIED	52645156, 52646365, 52645080, 35696052,
		(X83413) U88 [Human herpesvirus 6]			33656970, 52646317, 33657084, 265017.
					21906768, 21906769, 35695917, 33657109,
					52645129, 33657182, 27486261, 27486262, 33657349, 27486265, 18108387
2248	95412996 (4495, 4496)	1	Contains protein domain (PF00089) - cathepsin		264488, 264259, 264907, 29331830, 264909.
		02/refINP_004123.1/pHABP - hyaluronan-binding	Trypsin		265007, 265009, 264595, 21906754.
		protein 2			65274444, 264603, 265019, 264762, 264448,
	•				264288, 264689, 21906766, 55811957,
					265021, 264691, 18108374, 264634, 264635,
,					264636, 264555, 264638, 264557, 264558,
					264559, 18108383, 83373044, 18108385,
2249	94685662 (4497, 4498)	94685662 (4497, 4498) Novel Protein sim GRank nitangaast (45107772) - TCST11	nij4038461 (AE107772) - TCST(1   Costains protein domain (BE00616)		264758 264638 264637
		[Trypanosoma cruzi]	TPR Domain		704100, 204020, 204030, 204031
2250	79827508 (4499, 4500)	79827508 (4499, 4500) Novel Protein sim. GBank gij3738140jembjCAA212411 -		UNCLASSIFIED	264908, 18108374
		(AL031652) valyl-trna synthetase, mitochondrial precursor (Schizosaccharomyces pombe)			
2251	87385863 (4501, 4502)	87385863 (4501, 4502) Novel Protein sim. GBank gij3218467jemb CAA07090.11 -		UNCLASSIFIED	264259, 35696052, 264508, 56182435,
		(AJ006529) putative phosphatase [Gallus gallus]			265009, 264592, 264593, 264760, 264448,
					264684, 264288, 264690, 264628, 55811576,
					264555, 264556, 264557, 284558, 264559,
252	87735867 (4503, 4504)		Contains protein domain (PF01813) - synthase		264092, 264094, 264259, 29331822.
		AF14531 - (AF145316) vacuolar	ATP synthase subunit D		66714117, 29331828, 264102, 264103.
		proton pump delta polypeptide [Homo sapiens]			264104, 264105, 264109, 284112, 264511,
	•				265007, 60433356, 265010, 18108351,
					21906767, 21906768, 264691, 263974,
2253	91010703 (4505, 4506)			T	203317, 204400, 204307
7	7222 1227 1227			UNCLASSIFIED	652/45/Z, Z65U19

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18108396, 65274572, 56182575, 22278997, 22278998, 264259, 29331822, 29331827, 264905, 66712502, 264908, 264909, 264905, 2646905, 6712502, 264908, 60432229, 33657064, 87168559, 18108351, 26448, 264683, 264589, 264699, 56181562, 265021, 60170615, 264690, 33657109, 60431528, 18108374, 52644332, 56182323, 18108385, 22278000, 22278000, 26482	56182575, 265020, 264905, 264906, 264908, 35696423, 264511, 264635, 55812038, 264758, 265018, 265019, 264605, 264760, 264583		56182575, 22278999, 29331822, 29331825, 60432289, 29331827, 35696052, 264508, 66712502, 52644045, 56182435, 265006, 265008, 265009, 60433356, 55812038, 265010, 265017, 265019, 264288, 284389, 21906765, 21906767, 55811957, 35695817, 5264150, 33657109, 55811576, 64387374, 64487373	22278994, 22278997, 264907, 264828, 52644150, 18108381, 264693, 18108374	264686, 264488, 264768, 264769, 264691, 264508, 264905, 264509, 264906, 264907, 264908, 264908, 264908, 264908, 264908, 264510, 264511, 264512, 265007, 265009, 264638, 264639, 264757, 264758, 18108385, 265011, 264760, 264564, 264565, 264764, 264566, 264686, 264466	264689, 264910, 264764	22278995, 22278997, 22278999, 264259, 265006, 265007, 265009, 60433438, 21906754, 265010, 265011, 265017, 264448, 264683, 264689, 21906765, 21906768, 35695917, 265021, 18108374, 264638, 22279000, 22279002, 264568,
UNCLASSIFIED	oxidase	UNCLASSIFIED			UNCLASSIFIED	UNCLASSIFIED	ribosomalprot
	Contains protein domain (PF00090) - oxidase Thrombospondin type 1 domain					Contains protein domain (PF00096) - UNCLASSIFIED Zinc finger, C2H2 type	Contains protein domain (PF01305) - ribosomalprot Ribosomat protein L15 amino terminal region
2259   95364155 (4517, 4518) Novel Protein sim. GBank gil4884140 emb CAB43278.1  - (AL.050110) hypothelical protein [Homo sapiens]	88084119 (4519, 4520) Novel Protein sim. GBank gij3080663 (AC004614) - similar to f-spondin proteins AB006086 (PID:g2529225) [Homo sapiens]	88074157 (4521, 4522) Novel Protein sim. GBank gij3334526jembjCAA16138j - (AL021306) predicted using FGENEH [Homo saplens]		87602495 (4525, 4526) Novel Protein sim. GBank gij3341697 (AC003672) - hypothetical protein [Arabidopsis thaliana]	Novel Protein sim. GBank gil1657601 (U66220) - unknown  Nannocysiis exedens	86918663 (4529, 4530) Novel Protein sim. GBank gil477072 pir  A48018 - mucin 7 precursor, salivary - human	87773458 (4531, 4532) Novel Protein sim. GBank gil3150479 (AF067212) - partial CDS [Caenorhabditls elegans]
.59 95364155 (4517, 451		_					2266   87773458 (4531, 453

2267	87395838 (4533, 4534)	87395838 (4533, 4534) Novel Protein sim. GBank gij3560229jemb CAA20697.1 -		UNCLASSIFIED	35696286, 264259, 29331824, 29331825,
		(Accounts) hypometical protein (actizosaccuaromyces, pombe)	*		35696052, 29331828, 264905, 264509, 264907, 264908, 264909, 264512, 265009
			-		264910, 264593, 33657402, 265010, 265018,
					264762, 264448, 264288, 264369, 264768,
					52644229, 35695917, 264691, 33657023.
					18108382, 33657109, 35698423, 264634,
2268	95603867 (4626 4636)	The state of the s			18108381, 87168518, 264566
2007	(4323, 4330)	opospor (4555, 4550) ivovel Protein Sim. GBank		cadherin	264488, 264259, 264509, 264595, 265010,
		(B) 728832 SP P39189 ALUZ_HUMAN - !!!! ALU SUBFAMILY  SR WARNING ENTRY !!!			265017, 264766, 18108385, 264486
2269	88177977 (4537, 4538)	88177977 (4537, 4538) Novel Protein sim GBank oil102418loiri1917885 - T-037		INCI ACCIETED	56183676 60433040 366007 366000
_	(222)	motoria force anni: Openia giptora loppin por recording anni openia		UNCLASSIFIED	301623/3, 6043Z049, Z630U/, Z630U9,
		protein - itali iiy (Orosophila melanogaster)			264591, 87168559, 264805, 18108351,
					21906764, 265020, 264629, 60431528.
_	80410327 (4530 4540)				204036, 18106365, 18108387, 60432113
277	004 10327 (4539, 4540)				264763
	91010392 (4541, 4542)			cyto450	264909, 56182435, 265008, 55812038,
					55811957, 33657023, 264693, 33657109.
					55810764, 55811576, 56182323
7/77	84208220 (4543, 4544)			UNCLASSIFIED	264905, 264908
	95014271 (4545, 4546)	95014271 (4545, 4546) Novel Protein sim. GBank gil4176370 (AC005058) - similar			52645156, 22278996, 22278999, 60432049,
			Glutaredoxin		264259, 29331822, 29331824, 29331825,
		AC004392 (PID:g3367519) [Homo sapiens]			29331826, 29331827, 35696052, 264909.
					265006, 264593, 60433438, 21906754,
					265018, 264689, 21906765, 21906766,
					21906767, 21906769, 265021, 265022,
		`			60170615, 264691, 33657023, 264693,
				•	33657109, 27486264, 18108376, 35696423.
					35695855, 264630, 52644332, 264558,
27.74	01640217 /4647 46401				56182323, 22279002
	91040217 (4347, 4340)	olovoz II (4347, 4340) Novel Protein Sim. GBank gij 1480112jembjCAA67961	Contains protein domain (PF00538) - histone	histone	52645156, 22278997, 22278999, 52645080,
		(ASSO42) HF1-BF/4 protein [Mus musculus]	linker histone H1 and H5 family		29331824, 29331825, 29331826, 29331827,
					29331828, 264905, 264908, 52644045,
					264511, 265008, 265009, 60170831, 264591,
					21906754, 33109954, 265011, 265018,
,					18108351, 264448, 264288, 264684, 264766.
					21906765, 21906766, 21906767, 21906768.
					52644150, 264693, 18108384, 35695763,
					18108374, 35696423, 264634, 264557.
					284638, 52644332, 83373044, 18108385,
3766	00000 00000				56526486, 87168518, 22279002
6773	66062301 (4348, 4550)	odoczou (4549, 4550) Novel Protein sim. GBank gij3165406 (AC004755) - [fos37502_2 [Homo sapiens]	Contains protein domain (PF00122) - Iransport E1-E2 ATPase	transport	
2276	11287447 (4551, 4552)			INCI ASSIEIED	DRAKER DRAKER
				מוטביים וויה	204333, 204333

2277	88084123 (4553, 4554)	88084123 (4553, 4554) Novel Protein sim GBank nij2880070 (ACCOLATE)			
		to murine leucine-rich repeat protein; possible role in neural	Brizodova (Acodo 144) - Similar   Contains protein domain (PF00560) - glycoprotein at protein; possible role in neural (Leucine Rich Repeat	glycoprotein	22278999, 35696052, 265008, 265019,
		development by protein-protein interactions; 93% similarity to D49802 (PID:o1369906) Homo kanienal			204309, 253020, 253022, 55810764, 264404.   22278002
2278		Novel Protein sim. GBank gil2618702 (ACO02510)			
		unknown protein [Arabidopsis thaliana]	-	UNCLASSIFIED	18108394, 22278997, 28331826, 60433356, 60433438, 21906754, 265018, 33657023,
2279		80419375 (4557, 4558) Novel Protein sim. GBank			264639, 83373044, 264565
		gij19714 spip13983 EXTL_TOBAC - EXTENSIN PRECURSOR (CELL WALL HYDROXYPROLINE-RICH GLYCOPROTFIN)		UNCLASSIFIED	264768, 264565
2280					
2281		95293048 (4561, 4562) Novel Protein sim GBank oil 4240200 della 6 2 402 8 11			265008, 33109954, 265010, 265019, 265020
			Contains protein domain (PF00400) - Itransport	transport	264092, 264259, 29331822, 29331824,
			. O-Dela lepeal		29331826, 35696052, 264107, 264906,
					204909, 5264045, 265006, 33657402,
_					60433356, 264758, 265011, 265019, 264681,
					264663, 264684, 264686, 21906765,
	•				21906767, 21906768, 21906769, 60170615,
					264690, 52644150, 18108362, 264692.
					18108368, 18108374, 263978, 264631,
					18108381, 264559, 18108385, 56526486,
2282	87602829 (4563, 4564)	87602829 (4563, 4564) Novel Protein sim. GBank gil1537070 (1163840)			22279000, 264566, 264567
		nucleoporin p54 [Rattus norvegicus]		UNCLASSIFIED	264488, 264259, 29331822, 29331824,
					29331827, 29331828, 29331830, 33657402,
					60433438, 87168474, 265019, 18108351,
			•		21906767, 21906769, 55811957, 33657023,
_					52645129, 33657109, 33657182, 27486262,
2283	95362386 (4565, 4566)	95362386 (4565, 4568) Novel Protein sim. GBank		T	263972, 55811576, 87168518, 20281169
		gil2495729lsplQ92556lY281_HUMAN - HYPOTHETICAL	-	UNCLASSIFIED	60424179, 56182575, 22278994, 35696286,
		PROTEIN KIAA0281 (HA6725)			22278997, 22278999, 29331822, 29331824,
			-		56182181, 29331825, 29331827, 35696052,
•					29146499, 264905, 86712502, 264908,
					265007, 265009, 60432229, 264593,
					60431735, 60433356, 33109954, 33657084,
					55811386, 87168474, 265010, 265011,
					265018, 265019, 55811150, 264683, 264369.
				<del></del>	264288, 264688, 21906765, 21908767,
					21906768, 29148627, 21906769, 55811957,
				<u> </u>	265020, 265022, 33657182, 27486261,
					18108370, 264628, 18108374, 55810764.
					18108379, 55811576, 35696423, 35695855,
					264630, 60431850, 263981, 18108382,
					83373044, 18108385, 18108387, 60432113,
					222/8000 2844B2 2645B7

60424179, 52644507, 18108394, 52646842, 22278994, 35686286, 22278993, 22278999, 284259, 80432049, 29331822, 29331824, 29331824, 29331825, 60432289, 29331822, 29331827, 35686052, 29331830, 5264045, 56182435, 33657402, 60433438, 33109954, 21908754, 85858542, 87186559, 265018, 265019, 55811150, 284682, 264369, 21908765, 21908766, 21908767, 21908769, 21908769, 21908769, 21908769, 21908769, 21908769, 21908769, 21908769, 21908769, 21908769, 21908769, 21908769, 21908769, 21908769, 33657782, 255018, 25510764, 35695855, 60431850, 87168518, 87168513, 35695855, 60431850, 87168518, 60437113, 2646482, 27486265, 18108376, 25810764, 35696823, 35695855, 60431850, 87168518, 60431850, 87168518, 60431850, 87168518, 60431850, 87168518, 60431850, 8716850, 8716851850, 8716851850, 8716851850, 8716851850, 87168518650, 8716851850, 8716851850, 8716851850, 8716851850, 8716851850, 8716851850, 8716851850, 8716850	collagen 35696052, 284905, 264907, 264909, 264509, 264509, 264760, 18108351, 264682, 264763, 284685, 264768, 264693, 264629, 35695855, 264634, 264634, 264635, 264634, 264644,	kinase 35696286, 56182435, 60170831, 264591, 60432229, 264592, 264593, 264594, 264595, 55812038, 264596, 87168474, 35695917, 264692, 55811576, 264555, 264557	391) - UNCLASSIFIED	1m7 22278995, 56994075, 22278997, 22278999, 264269, 264269, 264269, 265017, 265018, 265019, 18108351, 264448, 21906765, 21906766, 21906767, 21906768, 33657023, 264557, 22279000, 22279000	F00400) - kinase	Contains protein domain (PF00400) - UNCLASSIFIED 264683 WD domain, G-beta repeat	struct 18108394, 264907, 265006, 265009, 33109954, 52646317, 265010, 18108351, 26263 26263 26263
Novel Protein sim. GBank gilz498797jsp G64311jPNaD_MOUSE - PROTEIN N. TERMINAL ASPARAGINE AMIDOHYDROLASE (PROTEIN NH2-TERMINAL ASPARAGINE DEAMIDASE) (NTN- ASPARAGINE AMIDOHYDROLASE) (PNAA) ASPARAGINE AMIDOHYDROLASE) (PNAA)	Novel Protein sim. GBank gij3342234 (U93309) - nuclear antigen EBNA-1 (Cercopithecine herpesvirus 15]	Novel Protein sim. GBank gij3873414 (U00043) - similar to D. melanogaster trithorax protein [Caenorhabditis elegans]	gil630905[pirt]S42731 - collagen (Hemicentrotus pulcherrimus)	2_HUMAN - !!!! ALU SUBFAMILY	gi 2887497 (AC004144) - 	1	gi 3253120 (AC005175) - ]
ASPA 95414955 (4567, 4568) Novel Protein sim. GBank gilz498797[sp[064311]PN TERMINAL ASPARAGINE NH2-TERMINAL ASPARAGINE AMIDASE) (PNAD) ASPARAGINE AMIDOHY	2285   87781484 (4569, 4570)   Novel Protein sim. GBank antigen EBNA-1 (Cercopti	87737825 (4571, 4572)		94133083 (4575, 4576) '			2291   84295281 (4581, 4582)

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56182575, 35596286, 56994075, 29331824, 29331825, 35696052, 56182435, 60433438, 55812038, 33109954, 87168474, 87168559, 265018, 18108351, 264763, 264448, 284369, 264288, 56181562, 284769, 21908765, 21908765, 21908765, 21908762, 23657023, 264693, 65274620, 33657109, 27486264, 264629, 55810764, 55816768, 35695855, 56182323, 56528486, 87168518, 325279000, 284567	264488, 18108397, 35696286, 264092, 264259, 29331822, 29331826, 264906, 264908, 264511, 264512, 265009, 264910, 18108351, 264764, 264369, 264288, 264686, 265020, 265022, 264534, 35696423, 264631, 264637, 18108381, 56182323, 264530, 18108385, 564464, 264667, 264657, 264667, 26467,	22278997, 22278998, 22278999, 264259, 29331822, 60432289, 29331828, 3569052, 265018, 264684, 264289, 294886, 21906765, 21906766, 21906767, 21906768, 21906769, 265020, 265021, 264692, 33657199, 18108376, 35696423, 35695855, 264684, 22279000, 22279002, 264563, 264486	22278996, 6043289, 264682, 264683, 22278996, 6043289, 264682, 264683, 26689, 18108374, 263974, 263978, 26488, 65274572, 56182575, 22278997, 22278999, 264259, 29331822, 29331824, 29331826, 29331822, 29331824, 29331826, 29331826, 264908, 5264045, 56182435, 264112, 265006, 265007, 264910, 265009, 60433356, 33657402, 264595, 55812038, 21906764, 264786, 21906765, 21906767, 21906768, 21906769, 55811957, 265200, 265021, 264593, 33657109, 55811576, 56182323, 60170394, 83373044, 18108385, 55526486, 264640, 264885, 264886, 264886, 264881, 26528486, 264881, 26488
UNCLASSIFIED	sind	nuclease	UNCLASSIFIED  Iranscriptfactor
	Contains protein domain (PF00560) - struct Leucine Rich Repeat	Contains protein domain (PF00929) - nuclease Exonuclease	
Novel Protein sim. GBank gil4803672 emb CAB42643.1	87759213 (4585, 4586) Novel Protein sim. GBank gij3252981 (AF068921) - Ras- binding protein SUR-8 [Mus musculus]	86693580 (4587, 4588) Novel Protein sim. GBank gil2062680 (U88964) - HEM45 [Homo sapiens]	95312200 (4589, 4590) 80030781 (4591, 4592) 94321251 (4593, 4594) Novel Protein sim. GBank gil5689501 dbj BAA83034.1 - (AB028005) KIAA1082 protein [Homo sapiens]
2292   94328634 (4583, 4584)   Novel Protein sim. GBank (AJ133769) nuclear trans;			
229:	2283	2294	2295 2296 2297

60424179, 56181686, 22278995, 35696286, 22278999, 22278999, 2278999, 264490, 264256, 29331824, 66714117, 60424269, 35686052, 29331824, 66714117, 60424269, 35686052, 29331824, 66712502, 26182435, 264510, 265006, 6043343, 21009554, 264687, 18108351, 264682, 264689, 264689, 21806763, 264689, 21806763, 264689, 21806763, 264689, 21806763, 264689, 21806362, 3659517, 265020, 18108362, 35696423, 35695655, 264630, 264482, 264586, 60431850, 18108396, 56182323, 264558, 83373044, 18108396, 56182323, 22279000, 264482, 264567, 264482,	264369	264488, 22278998, 22278999, 264259,	[25551824, 00714117, 55050002, 204505, [264905, 264906, 264907, 264908, 264909.	265008, 284910, 265009, 264758, 265010,	87168559, 264600, 265018, 264760, 264762,	18108351, 264764, 264766, 264768, 264769,	21906766, 21906767, 35695917, 265021,	18108374 35505402 355053705, 10105370,	264636, 264638, 18108385, 22279002,	264563	264908, 264758, 265017, 21908765. 83373044, 264563	52644045, 265019, 264288, 33657023,	18108370, 18108385 264259 60432049 264907 264909 264910	60432229, 33857402, 265011, 265018.	264762, 264448, 264769, 264637, 264638.	264259, 29331824, 21906767, 33657182,	33657349	65274572, 22278996, 264908, 265006, 21906769, 264691, 264486
Collagen	UNCLASSIFIED							-			struct	UNCLASSIFIED	10,000	5		UNCLASSIFIED		
*													Cathaire action domain (DE00043)	Immunoglobulin domain				
Novel Protein sim. GBank gij3875051[emb CAB02849] - (Z81050) predicted using Genefinder; similar to collagen, cDNA EST EMBL:D65564 comes from this gene; cDNA EST EMBL:D69046 comes from this gene; cDNA EST yk366b12.3 comes from this gene; cDNA EST yk366b12.5 comes from this gene.		2300   94124346 (4599, 4600) Novel Protein sim. GBank gil244386 (AC002294) -	Ottatiown protein (Arabidopsis trailana)	•							91235725 (4601, 4602) Novel Protein sim. GBank gil2143637 pir  i84505 - calcium-  dependent actin-binding protein - rat	88084141 (4603, 4604) Novel Protein sim. GBank gil2887497 (AC004144) -	R34001 1 (Homo sapiens)	(AL049946) hypothetical protein [Homo sapiens]		94840434 (4607, 4608) Novel Protein sim. GBank	gi[2494162]sp[Q10005jVRY1_CAEEL - HYPOTHETICAL 39.9 KD PROTEIN T15H9.1 IN CHROMOSOME II PRECURSOR	90935911 (4609, 4610) Novel Protein sim. GBank gil4972686 gb AAD34738.1  - (AF132150) unknown [Drosophila melanogaster]
(281050) Novel Protein sim. GBank (281050) predicted using (281050) predicted using (CBNA EST EMBI:D65564 EST EMBI:D65564 EST EMBI:D6564 comes yk366b12.3 comes from this gene	2299  80193720 (4597, 4598)	0 94124346 (4599, 4600)								┪	2301   91235725 (4601, 4602) 	2302 88084141 (4603, 4604)	2303 94141439 (4605 4606)	7000.		2304 94840434 (4607, 4608)		2305 90935911 (4609, 4610)

	18108387, 264563, 264566, 264486		16.0030, 21.0030, 22.0030, 20.0031, 20.0031, 20.0031, 20.0031, 20.0031, 22.78999, 22.78999, 20.0031, 2	652/4620, 652/4/91		264512, 265007, 265008, 265009, 60433438, 33109954, 265010, 265011, 264603, 265017, 18108351, 264762, 2646831, 264288, 264369, 264688, 33657023, 20281149, 202811069, 264628, 2638125, 264628, 2658123, 264638, 18108385, 18108387, 87168518, 27272000, 264638
Kinasereceptor	CHISTOR LUNI	synthase	glycoprotein	INCI ASSIEIED	transcriptfactor	
Contains protein domain (PF00400) - kinasereceptor WD domain, G-beta repeat					Contains protein domain (PF00013) - transcriptfactor KH domain	·
2306   95334940 (4611, 4612)   Novel Protein sim. GBank   gil4929565 gb AAD34043.1 AF15180 - (AF151806) CGI-48   protein [Homo sapiens]	4614)	87608409 (4615, 4616) Novel Protein sim. GBank gil4758732[ref]NP_004522.1 pMOCS - molybdenum cofactor synthesis 2	Novel Protein sim. GBank gi[3878059]emb[CAB17070] - (299942) cDNA EST EMBL:D73444 comes from this gene; cDNA EST EMBL:D730905 comes from this gene; cDNA EST EMBL:D7208 comes from this gene; cDNA EST EMBL:D75030 comes from this gene; cDNA EST EMBL:D72944 comes from this gene; cDNA EST	4620)	87721189 (4621, 4622) Novel Protein sim. GBank gi[2137337 pir  148281 - gene mCBP protein - mouse	
95334940 (4611,						
230	2307	230	5308	2310	231	

WO 00/58473 PCT/US00/08621

(4625, 4626) NOI (AB27, 4628) NOI	2312 87549681 (4623, 4624)   Novel Protein sim. GBank gil2911264 (AC002550) - Unknown gene product [Homo sapiens]  2313 80042533 (4625, 4626)   Novel Protein sim. GBank gil3043626 db  BAA25477  - (AB011123) KIAA0551 protein [Homo sapiens]  2314 94313401 (4627, 4628)   Novel Protein sim. GBank gil3043626 db  BAA25477  -		INICI A SCIEICE	56182575, 56994075; 35696286, 22278996, 22278997, 22278999, 264269, 29331822, 29331824, 29331825, 29331826, 29331827, 25644045, 60432229, 60433356, 55812038, 33109954, 21906754, 87168474, 255018, 18108351, 264288, 52644229, 21906765, 21906765, 21906766, 21906767, 21906768, 21908769, 255022, 60170815, 33657023, 27486281, 27486264, 3569643, 35695855, 18108385, 22279000, 22279002
(AL 80430119 (4629, 4630)	Navel Frotein sim. Gbank gijoobb 7 4(emb CAB51401.1  - (AL035398) dJ796117.2 (CGI-51) [Homo sapiens]		UNCLASSIFIED	52644507, 52646365, 52646842, 22278996, 22278997, 22278999, 262259, 52645080, 29331822, 29331824, 60432289, 29331826, 29331827, 29331828, 35696052, 33656970, 52644045, 265008, 264593, 6043335, 264758, 33109954, 265010, 265017, 265019, 264298, 264369, 21906766, 21906766, 21906768, 35695917, 35644150, 33657023, 33657109, 52644129, 33659563, 18108374, 35696423, 35695855, 52644332, 22279000, 22279002, 264563, 264567
832) No (AF	94312191 (4631, 4632) Novel Protein sim. GBank gil5531827[gb AAD44488.1] - Contair (AF078856) p47 [Homo saplens]	Contains protein domain (PF00789) - glycoprotein		55811576, 264535, 56182223, 18100385, 52644507, 52645156, 52646365, 22278994, 22278995, 22278996, 22278996, 22278997, 22278998, 22278999, 264259, 29331822, 52645060, 29331824, 29331822, 5245600, 29331824, 29331825, 29331827, 35696052, 29331828, 264511, 265007, 265008, 265009, 26182435, 264511, 265007, 265008, 265009, 26182435, 264511, 265007, 265008, 265009, 265010, 81168559, 265017, 265017, 265019, 264681, 264763, 264448, 264689, 21906765, 21906764, 264689, 21906765, 21906765, 21906767, 21906768, 21906769, 35695017, 265021, 52644150, 33657023, 18108367, 18108376, 32696423, 35695855, 264631, 264566, 264669, 21908387, 8168318, 80432113, 22279000, 264566, 264657

2317	2317  87020571 (4633, 4634)			UNCIASSIFIED	22278008 60432040 264010 60432220
		`	·		264686, 264687, 264688, 264689, 264558,
3340	10001 3001 0000000				18108385
	(18939679 (4633, 4636)		1	UNCLASSIFIED	265006, 264910
ALS7	95101781 (4637, 4638)	93101761 (4637, 4638) Novel Protein sim. GBank gi[5262613]emb[CAB45746.1] -			264488, 264569, 18108396, 52646365,
		[suandes office in the interior of the interio			22278994, 22278995, 22278996, 56994075,
					33096266, 22278997, 22278998, 264259,
					52645080, 29331825, 29331826, 29331827,
					23331826, 2333183U, 36182433, 6U170831, 60433330 6043436, 33667403, 34006764
					50435E23, 60431733, 3363740E, E1806734, 52644306 R7168474 265044 R7168550
					265017, 265018, 265019, 18108351, 264448
					18108354, 264288, 264369, 52644229,
		•			21906764, 21906765, 21906766, 21906767,
				_	21906768, 21906769, 265021, 265022,
					52644150, 33657023, 52845129, 33657109.
					27486264, 33657349, 35695763, 18108370,
					18108376, 18108379,.35696423, 264558,
					83373044, 18108385, 56526486, 87168518,
					264564, 264565, 264566
2320	91622426 (4639, 4640)			kinase	22278994, 60432049, 60432289, 29331827,
		gij728837 sp P39194 ALU7_HUMAN - IIII ALU SUBFAMILY			264511, 265008, 52646317, 265017, 265019,
1		SO WARNING ENTRY !!!!			21906765, 18108372, 18108387, 22279002
2321	94320377 (4641, 4642)	94320377 (4641, 4642) Novel Protein sim. GBank gij3873837jemb CAB02700  -		UNCLASSIFIED	264488, 264687, 18108394, 264689,
		(281029) Similarity to S.pombe hypothetical protein			21906765, 18108397, 18108398, 21906767,
					21906768, 65274791, 22278995, 35695855,
_		from this gene; cDNA EST EMBL: T01062 comes from this			22278998, 265021, 265022, 264510, 265006.
	*	gene; cONA EST EMBL: T01321 comes from this gene;			264511, 264512, 265008, 60170615, 264555.
		CUNA EST EMBL: 102288 com			264636, 264556, 18108361, 264259,
	-				60432229, 33657023, 264557, 264558.
					264693, 60433356, 264559, 60433438,
					29331824, 18108365, 18108348, 18108384,
					29331825, 18108385, 33109954, 29331827,
					56526486, 29146499, 265011, 60432113,
					265017, 265018, 264508, 264563, 264482,
		,			264509, 18108351, 264448, 264907, 264682.
					18108370, 264683, 264908, 264288, 264909,
2322	87803165 (4643 4644)	T			18108334, 284488, 284387
	מיניים (דרטר) יייים ו	gijoo/gao/jembjCAbo1685.1j -   Procoopija mejanogasted	Contains protein domain (PF00105) - dehydrogenase	dehydrogenase	[22278996, 264907, 264511, 264757,
		Diosoprilla melanogasteri	snon chain denydrogenase		18108331, 264768, 264638

12	1840445 (4645, 4646)		Contains protein domain (PF00226) - eph DnaJ domain		22278994, 22278995, 22278997, 60432049, 264259, 29331822, 33658970, 264509, 56182435, 264511, 265008, 60433356, 60433438, 55812038, 33109954, 21906754, 8565824, 87168474, 265011, 81168559, 265017, 265019, 264760, 204681, 18108351, 264369, 264368, 18108355, 264687, 264688, 21906767, 21906768, 55811957, 35695917, 265021, 33657023, 18108362,
1963361	86633607 (4647 4648)				27486262, 55811576, 264631, 264555. 83373044, 87168518, 60432113, 22279002 264592, 264593, 265020
88165	074 (4649, 4650)	6377.1 -		associated	265020
84390	962 (4651, 4652)	84390962 (4651, 4652) Novel Protein sim. GBank gil231885 sp P29981 CP4C_BLADI - CYTOCHROME P450 4C1 (CYPIVC1)	Contains protein domain (PF00067) - cyto450 Cytochrome P450		265006, 264759, 35695855, 56182323
88081	648 (4653, 4654)	88081648 (4653, 4654) Novel Protein sim. GBank gil4240227Idbj BAA74892.1 -  (AB020676) KIAA0869 protein [Homo saplens]			
83388	428 (4655, 4656)	83388428 (4655, 4656) Novel Protein sim. GBank gij1245105 (U46463) - glutamine repeat protein-1 [Mus musculus]			264259, 264508, 264905, 264906, 264907, 264908, 264908, 265908, 265007, 264512, 264910, 264758, 265010, 264766, 264768, 264769, 33657023, 264693, 264628, 264631, 264634, 264638, 264688, 2646
8760	1478 (4657, 4658)	87604478 (4657, 4658) Novel Protein sim. GBank gil1169343 sp P42209 DIF6_MOUSE - DIFF6 PROTEIN	Contains protein domain (PF00735) - UNCLASSIFIED Cell division protein		60433438, 264595, 265017, 264766, 264692, 264629, 264635, 264638, 264638, 56182323, 60432113, 264566
8733	87335396 (4659, 4660)			UNCLASSIFIED	265017, 264685, 60432113, 264088
86990	<b>463 (4661, 4662</b>	86990463 (4661, 4662) Novel Protein sim. GBank gil5679136lgblAAD46874.1IAF16093 - (AF160934) BcDNA,LD14189 IDrosophila melanogaster)		transport	265009
8778	4182 (4663, 4664	2332 67784182 (4663, 4664) Novel Protein sim. GBank gi[2104452]emb[CAB08779] - (295397) unknown [Schizosaccharomyces pombe]		led	35696286, 22278998, 29331824, 60424269, 265006, 265008, 265018, 264448, 264764, 21906765, 35695917, 35695855, 264636, 22279000, 264566
8820	5958 (4665, 4666	88206958 (4665, 4666) Novel Protein sim. GBank gij38799985 emb]CAA92691.1 - [Z88318], CDNA EST CEMSD62F comes from this gene; cDNA EST EMBL:C07930 comes from this gene; cDNA EST EMBL:C09493 comes from this gene; cDNA EST yk415e8.3 comes from this gene; cDNA EST comes from this gene; cDNA EST		UNCLASSIFIED	56182575, 56994075, 29331826, 29331828, 264107, 33657402, 87168559, 264683, 35695917, 265021, 33657023, 263976
9431	9788 (4667, 466£	94319788 (4667, 4668) Novel Protein sim. GBank gil4986270lgblAAB52261.21-(U97002) similar to acyl-CoA dehydrogenases and epoxide hydrolases; Pfam domain PF00441 (Acyl-CoA_dh). Score=57.4, E-value=1.7e-16, N=2; contains similarity to Pfam domain PF00702 (Hydrolase), Score=57.4, E-value=16-13, N=1 [C	Contains protein domain (PF00441) - dehydrogenase Acyl-CoA dehydrogenase	dehydrogenase	<del>5</del> 6182575, 29331825, 21906768, 264636, 83373044

2335		80046103 (4669, 4670) Novel Protein sim. GBank gij3283350 (AF062378) -  calmodulin-binding protein SHA1 [Mus musculus]	Contains protein domain (PF00612) - struct IQ calmodulin-binding motif	struct	18108351, 21906769, 264555
2336		Novel Protein sim. GBank giļ1929056 emb CAA72805  - (Y12090) putative 3,4-dihydroxy-2-butanone kinase [Lycopersicon esculentum]		kinase	264907, 35695917, 18108379
2337		95345810 (4673, 4674) Novel Protein sim. GBank gi 4495063 emb CAB39181.1  - (Z85986) dJ108K11.3 (similar to yeast suppressor protein SRP40) [Homo sapiens]		UNCLASSIFIED	35696286, 22278999, 56182181, 29331825, 60424269, 56182435, 33657402, 55812038, 55811386, 265017, 265018, 265019, 21906768, 35695917, 264691, 33657023, 33657109, 263972, 35696423, 35695855, 60432113
2338		87634045 (4675, 4676) Novel Protein sim. GBank gij2224689 dbj BAA20829  - (AB002372) KIAA0374 [Homo sapiens]	Contains protein domain (PF00323) - UNCLASSIFIED Mammalian defensin	UNCLASSIFIED	18108394, 29331822, 66714117, 60433289, 284908, 56182435, 265009, 60433438, 264596, 285010, 265019, 18108354, 264288, 264389, 55811957, 265021, 33657023, 263976, 58811876, 264632, 56182323, 264639
2339		85663319 (4677, 4678) Novel Protein sim. GBank gij3873550jembjCAA22127j - (AL033534) serine-tich protein [Schizosaccharomyces pombe]		UNCLASSIFIED	35696286, 264592, 264369, 264691, 264558
2340					65274572, 22278994, 35696286, 22278997, 22278999, 264259, 29331822, 60432289, 29331826, 29331826, 265009, 33657402, 33109954, 265017, 265018, 264766, 264685, 21906769, 35698423, 87168518, 22779000
2341					264259, 264908, 264909, 264682, 22279000
2342		95334968 (4683, 4684) Novel Protein sim. GBank gij3874563jembjCAB02797j - (Z81042) similar to Yeast hypothetical protein YEY6 like; - CDNA EST yk206h5.3 comes from this gene; cDNA EST yk206h5.5 comes from this gene; cDNA EST comes from this gene; cDNA EST yk303h1.3 this gene; cDNA	Contains protein domain (PF00400) - kinase WD domain. G-beta repeat	kinase	264488, 65274572, 22278995, 22278996, 22278997, 22278998, 22278999, 60432049, 264259, 29331824, 29331824, 29331825, 66714117, 29331826, 60432289, 35696052, 264905, 264509, 29331830, 265006, 60170831, 6043229, 60433356, 87168474, 265017, 265019, 264418, 264369, 264288, 21906765, 21906766, 55811957, 35692917, 265020, 265022, 52644150, 33659217, 265020, 3569409, 18108370, 264568, 26468, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264568, 264
250		of (1944) (4069, 406b) Novel Protein sim. GBank gil4929741[gb]AAD34131.1[AF15189 - (AF151894) CGI-136 protein [Homo sapiens]		UNCLASSIFIED	264907, 264512, 265011, 264683
2344	79953198 (4687, 4688)			UNCLASSIFIED	264758
2345	94319799 (4689, 4690)	94319799 (4669, 4690) Novel Protein sim. GBank 9i 2506307 sp P13944 CA1C_CHICK - COLLAGEN ALPHA  1(XII) CHAIN PRECURSOR (FIBROCHIMERIN)	Contains protein domain (PF00092) - von Willebrand factor type A domain	collagen	264488, 264259, 66712502, 264759, 83373044, 264568

35696286, 22278998, 264259, 35696052, 29331828, 33657402, 60433356, 33109954, 87188559, 264603, 265019, 18108351, 264681, 264685, 21908768, 255021, 33657109, 55811576, 3589855, 264637, 52244332, 264557, 83373044, 22279000, 22279002	22278997, 264511, 264683, 264684, 264706, 264687, 264688, 264691, 264692, 55811576	18108394, 35696286, 264259, 35696052, 264508, 264508, 264907, 264908, 264909, 264907, 264908, 264909, 264907, 264908, 264909, 264909, 264909, 264909, 264909, 264910, 265007, 264591, 264593, 264593, 264594, 264757, 264762, 18108351, 264764, 264693, 264628, 264639, 35695917, 264693, 264628, 18108370, 264629, 18108374, 35699423, 264639, 83373044, 18108385, 264567, 264486	35696052, 29146499, 264909, 264369	22278998, 22278999, 264259, 29331822, 29331824, 29331825, 29331827, 29331828, 33109954, 21906754, 265010, 87168559, 265018, 265018, 284761, 264681, 264681, 264691, 264692, 35695855, 87168518, 22279000, 22279002, 264482	56182575, 264909, 265006, 264558	264259, 264448	29331826, 55812038, 265019, 264692, 264636	29331824, 264908, 265006, 265008
proteaseinhib		kinase		·	kinase	UNCLASSIFIED	dna_ma_bind	UNCLASSIFIED
Contains prolein domain (PF00515) - TPR Domain						Contains protein domain (PF00076) - UNCLASSIFIED RNA recognition motif. (a.k.a. RRM, IRBD, or RNP domain)	Contains protein domain (PF00076) - dna_ma_bind RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)	Contains protein domain (PF00076) - RNA recognition motif. (a.k.a. RRM, IRBD, or RNP domain)
94131820 (4691, 4692) Novel Protein sim. GBank gi[1255411 (U53153) - one short   Contains protein domain (PF00515) - proteaseinhib region of weak similarity to S. cerevisiae protease A Inhibitor TPR Domain 3 (SP:P01094) and another short region of weak similarity to S. cerevisiae glucose repression mediator protein (SP:P14922) [Caenorhabditis elegans]		95196133 (4695, 4696) Novel Protein sim. GBank gil1929056 emb CAA72805  - (Y12090) putative 3,4-dihydroxy-2-butanone kinase [Lycopersicon esculentum]	87776502 (4697, 4698) Novel Protein sim. GBank gil4884106 emb CAB43254.1 -  (AL050062) hypothetical protein [Homo sapiens]	<del></del>	88988042 (4701, 4702) Novel Protein sim. GBank gij728832 spjP39189 ALU2_HUMAN - !!!! ALU SUBFAMILY SB WARNING ENTRY !!!!	87337196 (4703, 4704) Novel Protein sim. GBank gil731637lsplP38760 YHH5_YEAST - HYPOTHETICAL RNA recognition motif 75.9 KD PROTEIN IN SPO13-ARG4 INTERGENIC REGION RBD, or RNP domain)	91638784 (4705, 4706) Novel Protein sim. GBank gil1346955 sp P48809 RB27_DROME - HETEROGENEOUS NUCLEAR RIBONUCLEOPROTEIN 27C (HNRNP 48) (HRP48.1)	87337199 (4707, 4708) Novel Protein slm. GBank gil7316371sp P38760 YHH5_YEAST - HYPOTHETICAL RNA recognition motif 75.9 KD PROTEIN IN SPO13-ARG4 INTERGENIC REGION RBD. or RNP domain)
H131820 (4691, 4692)	85330367 (4693, 4694)	95196133 (4695, 4696)	87776502 (4697, 4698)	88260594 (4699, 4700)	86968042 (4701, 4702)	87337196 (4703, 4704)	91638784 (4705, 4706)	87337199 (4707, 4708)
	2347	2348	_		2351			2354

2355	[91638786 (4709, 4710)]	91638786 (4709, 4710) [Novel Protein sim. GBank gil4938503lembiCAB43861.11.	Contains profess domain (DE00078)   Jan. Big.	Paid our	
		(AL078465) hnRNP-like protein (Arabidopsis thaliana)	RNA recognition motif. (a.k.a. RRM		30334073, ZZZ78999, 364360 30331836 30331836 30331830
			RBD, or RNP domain)		201253, 2331023, 2331020, 2331020.
					82148474 DEEDAD DEEDAT DOLGET
					2/1004/4, 2530/U, 2530/1/, 25458/,
					2 (augros, 2 (augrof, 21905/69, 264691,
2366	_				22279000
255		53327000 (4711, 4712) Novel Protein sim. GBank gij5138920 gb AAD40377.1  -			52644507, 22278995, 35696286, 22278996,
					22278997, 22278998, 22278999, 264259.
				•	29331824, 66714117, 29331825, 60432289,
					35696052, 29331828, 264908, 66712502,
					264512, 265007, 265008, 60170831,
_					60432229, 60433356, 60433438, 264758,
		-			52646317, 33109954, 21906754, 55811386, ~
					87168474, 265017, 265018, 264605, 265019,
					264681, 264682, 264448, 264369, 264288,
					264686, 264768, 21906765, 21906768,
					21906767, 21906768, 21906769, 265021,
					60170815, 33657109, 27486264, 35695763,
					55810764, 18108379, 35696423, 55811578
					35895855, 60170394, 56182323, 83373044.
					18108385, 56526488, 264404, 60432113
		-			22279000, 264482, 264563, 264566, 264486
2257	т				264567
3	_			UNCLASSIFIED	284488, 264769, 18108394, 264259
		914929741 gb AAD34131.1 AF15189 - (AF151894) CGI-136			29331822, 18108370, 18108374, 264510
		protein [Homo sapiens]			265017, 264482, 264563, 264762, 264565
2360	_				264566, 264369, 18108354
6007		91/1/0/0 (4/13, 4/15) Novel Protein sim. GBank gil4218005 (AC006135) - putative		UNCLASSIFIED	22278997, 22278999, 264509, 264905
_		Vicilin storage protein (globulin-like) [Arabidopsis thaliana]			264592, 18108351, 264681, 264682, 264769
				100	32833986, 18108374, 264556, 18108385,
2359	87755859 (4717, 4718)	87755859 (4717, 4718) Novel Protein sim. GBank gil 1086830 (1141264) . coded for			264482
		by C. elegans cDNA vk20f8 5: coded for by C. elegans		UNCCASSIFIED	35696286, 22278998, 264905, 264511,
		CDNA yk44q1,5; coded for by C. elegans cDNA yk12b7 5:			265007, 265008, 60433438, 264288, 264686,
		coded for by C. elegans cDNA vk36o6 5: coded for by C.			Z1906/09, Z050Z0, Z0469Z, 35695855,
		elegans cDNA vk20/8.5: coded for by C. elegans cDNA			Z04558, 565Z6486, Z64563
	$\neg$	yk16g12			
2360	_	in sim. GBank		UNCI ASSIFIED	22278997 29331828 263981 22279000
		(Z69904) cDNA EST yk428d5.3 comes from this gene:			00001777.
		cDNA EST yk428d5.5 comes from this gene			
		[Caenorhabditis elegans]			
230	94232191 (4721, 4722)	Novel Protein sim. GBank gi[746487 (U23514) - No			22278995, 22278999, 264512, 265009
		definition line found [Caenorhabditis elegans]			264757, 21906765, 65274620, 18108370,
		-			60431528, 18108374, 264635, 60170394,
					264482

			1		
7007	91121133 (4723, 4724)	91721193 (4723, 4724) Novel Protein sim. Chank   Protein Sim. Chank   Protein HEAVY   Qij1171093 sp P19706 MYSB ACACA - MYOSIN HEAVY		UNCLASSIFIED	66432289, 264539, 25331624, 25331624.
		CHAIN IB (MYOSIN HEAVY CHAIN IL)			60433356, 264448, 264682, 264683, 264369.
	٠				21906765, 21906768, 21906769, 60432113,
					22213000, 22213002
2363	95006635 (4725, 4726)	95006635 (4725, 4726) Novel Protein sim. GBank gi 854065 emb CAA58337  -  (X83413) U88 [Human herpesvirus 6]		UNCLASSIFIED	264907, 264629, 264635
2364	_		Contains protein domain (PF00389) - reductase	reductase	264488, 18108394, 264887, 18108398,
		gij5839830jgb AAD45886.1JAF14601 - (AF146018)	D-isomer specific 2-hydroxyacid		22278996, 56994075, 35696286, 22278997,
		hydroxypyruvate reductase [Homo sapiens]	dehydrogenases		22278998, 264259, 66714117, 29331825,
					35696052, 264509, 264905, 264906, 264907.
					264908, 66712502, 264909, 264511, 265006,
					264512, 265007, 265008, 33657402, 264758,
					21906754, 87168474, 265010, 87168559.
					264603, 265017, 265018, 265019, 264760,
					264762, 18108351, 264448, 264764, 264683,
					264684, 264288, 18108355, 264766,
				•	18108358, 264689, 18108359, 21906765,
			:		21906766, 21908767, 35695917, 265020,
					265021, 265022, 60170615, 52644150,
	ı,				264691, 33657023, 264692, 18108364,
	-			,	33657109, 18108368, 18108370, 18108374,
	-				35696423, 35695855, 264635, 264556.
					264557, 264639, 60170394, 83373044,
					18108383, 18108384, 18108385, 18108388,
					56526486, 264482, 264564, 264486
2365		94140746 (4729, 4730) Novel Protein sim. GBank gi[1840045 (U49082) -	٠	transport	22278996, 22278998, 22278999, 264907,
		transporter protein [Homo sapiens]			264909, 264910, 33657402, 264758, 264600,
					264766, 264687, 264689, 21906765,
					21906767, 21906768, 21906769, 265021,
				,	33657023, 33657109, 83373044, 264566
2366	94312388 (4731, 4732)			UNCLASSIFIED	52644507, 264259, 29331824, 29331825,
_					29331826, 29331827, 29331828, 264907,
					29331830, 264909, 264511, 265008,
					33657402, 264595, 52646317, 265017,
				•	265018, 265019, 264605, 264685, 264766,
					264689, 21906766, 21906769, 35695917,
					265020, 265021, 265022, 52644150,
					35695855, 52644332, 18108385, 18108387,
1	т		_		264564, 264566
2367		94140910 (4733, 4734) Novel Protein sim. GBank gi 1065457 (U40410) - C54G7.4		UNCLASSIFIED	35696286, 21906768, 55810764, 65274791,
2368	194727 4775 A77E	-	wu domain, G-beta repeat	f	204507
	_				264628

2260	104314334 16737 47381	2350 104344334 (4737 4730) [N   D   1   1   1   1			
	(1011, 1101)			Sunci	26443U7, 22646842, 33888288, 264092, 264094, 52645080, 35696052, 264107.
		saplens)			29331830, 52644045, 265006, 265007,
					265009, 52644298, 52644229, 264689.
					21906765, 21906768, 35695917, 265020,
					52644150, 263967, 33657109, 27486265,
					35695763, 18108370, 263974, 18108374,
			!		18108376, 52644332, 263981, 18108385
2370	(79804120 (4739, 4740)			UNCLASSIFIED	264508, 264909, 264596
2371					264369
2372	Į			UNCLASSIFIED	263967, 263981
2373	_	87418611 (4745, 4746) Novel Protein sim. GBank gil4589582 dbj BAA76813.1  -		Γ	29331826, 265010, 265019, 35695917,
	_	(AB023186) KIAA0969 protein [Homo saplens]			264634, 60432113
2374	_	94123665 (4747, 4748) Novel Protein sim. GBank gi[5105131 db] BAA80445.1   -	Contains protein domain (PF01138) - UNCLASSIFIED		265006, 265007, 265008, 265009, 265011,
		(AP000061) 246aa long hypothetical ribonuclease PH	3' exoribonuclease family		264766, 35695917, 35695855, 263981,
1		(Aeropyrum pernix)			264557, 264565
6/67		Novel Protein sim. GBank		UNCLASSIFIED	60432049, 29331824, 264907, 52644045,
		BILIDALI I JOHNAN JOSPAN BET MOUSE - SIGNAL			264512, 60433356, 21906/54, 52644296,
	•	(SB-BETA)			6/100339, Z04448, Z19U0/03, Z19U0/08,
		(2000)			Z1906/09, 3363/023, 16108368, 538113/6,   52644332
2376	_	87613744 (4751, 4752) Novel Protein sim. GBank gi 2645435 (AF007780) - CHD3	Contains protein domain (PF00628) - ATPase associated	ATPase associated 3	264259, 29331830, 264909, 264910, 265009,
		[Drosophila melanogaster]	PHD-finger		60433438, 21906754, 265017, 265018,
				<u> </u>	265019, 264682, 264288, 264685, 21906767,
					263972, 35695855, 87168518, 60432113
2377	95319689 (4753, 4754)	95319689 (4753, 4754) Novel Protein sim. GBank gij5257005 gb AAD41239.1  -	Contains protein domain (PF01388) - UNCLASSIFIED		18108394, 65274572, 22278997, 22278999,
		(AF083249) Rb binding protein homolog [Homo sapiens]	ARID DNA binding domain		264095, 29331822, 29147620, 29331824,
					66714117, 29331825, 29331826, 29331828,
_					33656970, 29146498, 29146499, 264509,
					265006, 265007, 265008, 265009, 60170831,
					265010, 265011, 265018, 55811150,
		•			18108351, 264764, 264288, 21906767,
					21906768, 29148627, 29148629, 265021,
					33857023, 33657109, 18108370, 18108374,
			•		18108379, 35696423, 264556, 83373044,
					18108385, 18108388, 56526486, 22279000.
2378		A CONTRACTOR OF THE CONTRACTOR			22279002, 264563
		34137032 (4733, 4730) Nover Protein Sim. GBank gi 10/2198 (U40942) - No		UNCLASSIFIED	65274572, 56182575, 35696286, 264259,
		definition line found [Caenomabditis elegans]			29331822, 29331824, 66714117, 29331825,
				<del></del>	29331826, 60432289, 29331827, 56182435,
		-			264510, 265009, 60433356, 87168474,
					265011, 265018, 264288, 21906765,
					33657023, 264557, 56182323, 83373044,
2370	65444324 (47E7 47ED)	1-100 min distant			18108385, 22279002, 264482
0		Novel Protein Sim. Gibank gri3337357 (AC004481) -   Invoothetical protein [Arabidoosis Ihaliana]	Contains protein domain (PF00400) - kinase		265017, 264288, 21906768
		formula made and the state of t	משלבו שיים מיים ובחתו איים ובחתו ו		

264908, 264910, 265011	56182575, 264091, 264093, 264259, 29331825, 264105, 264906, 60433356, 21906754, 265017, 265019, 264683, 264288, 264685, 264686, 264681, 264691, 264692, 264693, 55811576, 284636, 264567	29331824, 60432289, 264905, 264596, 21906754, 264769, 265022, 264693, 263967, 33657109, 264629, 264631, 264558, 83373044, 60432113, 264482	265009, 21906765, 21906766	264488, 52644507, 52645156, 52646365, 35696286, 22578999, 52645080, 29331824, 29331826, 25645080, 29331824, 29331826, 35696052, 29331828, 2564008, 29331824, 264628, 35696052, 29331828, 256908, 265001, 265011, 265017, 265018, 264639, 25244296, 265011, 265017, 265018, 264633, 264639, 26520, 5264150, 33657109, 33657182, 27486261, 27486262, 27486264, 33657109, 33657182, 27486265, 3569573, 18108374, 35695855, 264634, 264555, 264556, 264557, 52644312, 2248887, 22278999, 28278996, 25504045, 265006, 265008, 265009, 26500	26448, 264683, 264665, 264687, 264689, 21906765, 21906766, 21906767, 21906769, 25811957, 265021, 265022, 264690, 264692, 65274620, 33657109, 18108370, 264631, 5264332, 22278900, 22279002, 264563, 264565, 264567, 264267, 264267, 264267, 264267, 264267, 264267, 264267, 264267, 264682, 21906765, 21906769, 29148629, 36695917, 265621, 264690, 33657109, 264628, 18108376, 83373044, 60432113, 22279000, 264564, 264566, 264487
collagen	helicase	transport	glycoprotein	struct UNCLASSIFIED	glycoprotein
Contains protein domain (PF01410) - collagen Fibrillar collagen C-terminal domain	Contains protein domain (PF00385) - helicase 'chromo' (CHRromatin Organization MOdifier) domain	Contains protein domain (PF00628) - transport PHD-finger	Contains protein domain (PF00059) - glycoprotein Lectin C-type domain	Contains protein domain (PF00595) - struct PDZ domain (Also known as DHR or GLGF). UNG	
Novej Protein sim. GBank gij4502939jrefjNP_001845.1pCOL1 - collagen, type XI, alpha 1	87608241 (4761, 4762) Novel Protein slm. GBank gil4455609 emb CAB36555  - (AL031846) dJ742C19.5 (novel Chromobox protein) (Homo sapiens)	91225982 (4763, 4764) Novel Protein sim. GBank gil4325130 gb AAD17276  - (AF119716) dMi-2 protein [Drosophila melanogaster]	87442841 (4765, 4766) Novel Protein sim. GBank gil1902982 dbj BAA19005  - (D89049) lectin-like oxidized LDL receptor [Bos taunus]	95354766 (4767, 4768) Novel Protein sim. GBank gil2462851 (AF016252) - Spinophilin [Rattus norvegicus]	94742649 (4771, 4772) Novel Prolein sim. GBank gil4929699igb AAD34110.1 AF15187 - (AF151873) CGI-115 protein [Homo sepiens]
	87608241 (4761, 4762)	91225982 (4763, 4764)	87442841 (4765, 4766)		94742649 (4771, 4772)
2380	2381	2382	2383	2385	2386

2386 11424604 (4775, 4776) 2389 95310650 (4777, 4778) Novel Protein sim. GBank gild758058jrefilnY_004372.1pCREB - CAMP responsive element binding protein-like 1 (D84103) mitochondrial DNA polymerase gamma [Homo saplens] 2380 94320912 (4778, 4780) Novel Protein sim. GBank gild64239jdbjjBAA12223j - (D84103) mitochondrial DNA polymerase gamma [Homo saplens] 2381 80036194 (4781, 4782) 2382 94245016 (4783, 4784) Novel Protein sim. GBank gild240169jdbjjBAA74863.11-				total
95310850 (477, 4778) Novel Protein sim. GBank gli4758058[ref]NP_004372.element binding protein-like [94320912 (4779, 4780) Novel Protein sim. GBank (D84103) mitochondrial DN saplens]  80036194 (4781, 4782) Novel Protein sim. GBank (B425016 (4783, 4784) Novel Protein sim. GBank			UNCLASSIFIED	264595
gil4758058jrefjNP_004372 element binding protein-like 94320912 (4779, 4780) Novel Protein sim. GBank g (D84103) mitochondrial DN saplens] 80036194 (4781, 4782) Novel Protein sim. GBank		Contains protein domain (PF00170) - dna ma bind	dna rna bind	284488, 22278998, 22278999, 264509,
Selement binding protein-like   Selement binding protein sim. GBank   Selement binding protein sim. GBank   Selement binding protein sim. GBank   Selement binding protein-like   Selement binding   Selement bind	1 pckeb - caMP responsive	bZiP transcription factor	1	264905, 264906, 264907, 264908, 264909,
94320912 (4779, 4780) Novel Protein sim. GBank (D84103) mitochondrial DN saplens] 80036194 (4781, 4782) 84245016 (4783, 4784) Novel Protein sim. GBank	-			265006, 264511, 264512, 264910, 264591,
94320912 (4779, 4780) Novel Protein sim. GBank (D84103) mitochondrial DN saplens] 80036194 (4781, 4782) Novel Protein sim. GBank				21908754, 264601, 264604, 264761,
94320912 (4779, 4780) Novel Protein sim. GBank (D84103) mitochondrial DN sapiens] 80036194 (4781, 4782) 84245016 (4783, 4784) Novel Protein sim. GBank				18108351, 264764, 264288, 264766, 264768,
94320912 (4779, 4780) Novel Protein sim. GBank (D84103) mitochondrial DN saplens] saplens   80036194 (4781, 4782)   84245016 (4781, 4782)   Novel Protein sim. GBank				264769, 21906765, 21906768, 264692,
94320912 (4779, 4780) Novel Protein sim. GBank (D84103) mitochondrial DN sapiens] 80036194 (4781, 4782) 84245016 (4783, 4784) Novel Protein sim. GBank				264693, 35696423, 264635, 264636, 264555.
94320912 (4779, 4780) Novel Protein sim. GBank (D84103) mitochondrial DN saplens  80036194 (4781, 4782) 84245016 (4783, 4784) Novel Protein sim. GBank				83373044, 22279000, 264486
(D84103) mitochondrial DN saplens  80036194 (4781, 4782) 84245016 (4783, 4784) Novel Protein sim. GBank	ji 1644239 dbj BAA12223  -	Contains protein domain (PF00476) - polymerase	polymerase	52644507, 56182575, 22278995, 35696286,
80036194 (4781, 4782) 84245016 (4783, 4784) Novel Protein sim. GBank	lA polymerase gamma (Homo	DNA polymerase family A		22278996, 22278997, 22278999, 29331822,
80036194 (4781, 4782) 84245016 (4783, 4784) Novel Protein sim. GBank				29331825, 29331826, 35696052, 264905,
80036194 (4781, 4782) 84245016 (4783, 4784) Novel Protein sim. GBank				52644045, 265009, 264758, 264759,
80036194 (4781, 4782) 84245016 (4783, 4784) Novel Protein sim. GBank				33109954, 52644296, 85658542, 265011,
80036194 (4781, 4782) 84245016 (4783, 4784) Novel Protein sim. GBank				265017, 265018, 264605, 52644229,
80036194 (4781, 4782) 94245016 (4783, 4784) Novel Protein sim. GBank				21906765, 21906767, 21906768, 21906769.
80036194 (4781, 4782) 94245016 (4783, 4784) Novel Protein sim. GBank				35695917, 52644150, 33657023, 33657109.
80036194 (4781, 4782) 94245016 (4783, 4784) Novel Protein sim. GBank				33657349, 35695763, 18108370, 18108374,
80036194 (4781, 4782) 94245016 (4783, 4784) Novel Protein sim. GBank				18108376, 35696423, 35695855, 264555,
80036194 (4781, 4782) 94245016 (4783, 4784) Novel Protein sim. GBank				52644332, 56182323, 60170394, 83373044,
80035194 (4781, 4782) 94245016 (4783, 4784) Novel Protein sim. GBank				56526486
94245016 (4783, 4784) Novel Protein sim. GBank			UNCLASSIFIED	263976
	gi 4240169 dbj BAA74863.1  -	Contains protein domain (PF00560) - nuclease	nuclease	35696286, 35696052, 264508, 264905,
(AB020647) KIAA0840 protein (Homo sapiens)	ein [Homo sapiens]	Leucine Rich Repeat		264509, 264906, 264907, 264908, 264909,
		,		264510, 264511, 264512, 264910, 265009.
				264591, 264758, 264600, 264604, 264762,
				264448, 264764, 264369, 264766, 264768,
				264769, 264689, 35695917, 264629,
				18108374, 263978, 35696423, 35695855,
				264631, 264634, 264635, 264636, 264637,
				264638, 60170394, 264639, 264565, 264486

- 7 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0		dna_ma_bind   264259, 29331824, 264910, 264265, co3u21,   83373044, 18108387, 264563, 264566
		m_eup
Ribosomal protein L10	Contains protein domain (PF00096) - dna_ma_bind Zinc finger, C2H2 type	
	94323268 (4787, 4788) Novel Protein sim. GBank gil4159888 (AC004908) - zinc finger protein from gene of uncertain exon structure; similar to Q99676 (PID:g3025333) [Homo sapiens]	95287212 (4789, 4790) Novel Protein slm. GBank glj5712756[gbJAAD47638.1 AF16079 - (AF160798) calcium
E COOL COOL COOL COOL COOL COOL COOL COO	94323266 (4787, 4788)	95287212 (4789, 4790)

52246365, 18108397, 56182575, 35696286, 22278997, 22278999, 60432049, 284259, 29331822, 29331824, 29331825, 29331826, 29331827, 29331828, 264106, 264907, 29331830, 66712502, 264110, 60170831, 264591, 33657402, 60433438, 55812038, 33109954, 21508754, 33657084, 8746842, 26428, 264766, 2644229, 2190676, 265020, 265021, 60170615, 264692, 33657023, 65274620, 52645129, 2365702, 18108388, 87168518, 22279002, 264482	52844507, 52645156, 56182575, 264259, 29147620, 264905, 264907, 264908, 264909, 264910, 264758, 52644296, 264603, 264604, 264762, 264681, 264764, 18108357, 264769, 21906768, 264693, 264628, 264635, 264638, 264639, 264564		56994075, 22278997, 22278998, 22278999, 264259, 264259, 29331822, 66714117, 60432289, 264906, 29331820, 56182435, 284112, 264910, 33109954, 21906754, 264600, 265017, 265018, 265019, 264764, 264765, 21906765, 21906766, 265020, 265022, 60170615, 33655027, 265020, 265022, 60170615, 33657023, 18108370, 264556, 60170394, 264556, 264564, 264566, 264687	21806766, 52646842, 56994075, 33657182, 27486262, 52644296, 265017
nuclease		UNCLASSIFIED	UNCLASSIFIED	phosphatase .
Contains protein domain (PF00560) - nuclease Leucine Rich Repeat				
2396   95096700 (4791, 4792) Novel Protein sim. GBank gil106322lpir  B34087 - hypothetical protein (L1H 3' region) - human		2398   88047689 (4795, 4796) Novel Protein sim. GBank gi 3258609 (AC005178) -   H53_GS1 (Homo sapiens)	) Novel Protein sim. GBank gij786117 (L41834) - nuclear protein [Ensis minor]	91214116 (4799, 4800) Novel Protein sim. GBank gil2352822lgbjAAB69285.11- (AF008945) glucose-6-phosphatase [Haplochromis nubilus]
95096700 (4791, 4792)		88047689 (4795, 4796	87738965 (4787, 4798	
2396	2397	2398	7338	2400

2401	91214118 (480)	31, 4802)	2401   91214118 (4801, 4802)   Novel Protein sim. GBank gil2352822[gbJAA869285.1] - (AF008845) glucose-8-phosphatase [Haplochromis nubilus]		phosphatase	52644507, 52645156, 52644229, 264688, 21906764, 21906765, 52646395, 52646842, 21906764, 21906765, 52646395, 52646842, 21906768, 21906768, 21906768, 2278995, 35695917, 56994075, 35696286, 22278996, 22278997, 265020, 22278998, 22278999, 264259, 33657023, 52645080, 264693, 23557182, 29331827, 3696952, 27486261, 27486262, 33659762, 264106, 264905, 35696423, 3569565, 264106, 264905, 35696423, 35696855, 265006, 265007, 265008, 265009, 264637, 52644296, 87168474, 265010, 87168559, 60432113, 265017, 265018, 265019, 264563, 264563, 264288
2402		03, 4804)	91221408 (4803, 4804) Novel Protein sim. GBank gi 4689258 gb AAD27832.1 AF12185 - (AF121859) sorting nexin 9 [Homo sapiens]			264907, 264908, 264909, 264566
2403		05, 4806)	JAF15181 - (AF151811) CGI-53	Contains protein domain (PF00062) - C-type lysozyme/alpha-lactalbumin family		22276999, 35696052, 265018, 264686, 264693, 83373044, 264567
2404		07, 4808)	gij2315796 (AF016685) - similar genases [Caenorhabditis	Contains protein domain (PF00106) - dehydrogenase short chain dehydrogenase	dehydrogenase	35696286, 29331826, 35696052, 265008, 265018, 21906769, 264564
2405		09, 4810			UNCLASSIFIED	35696286, 29331822, 265007, 21906754, 265017, 265018, 265019, 264763, 264369, 21906765, 35695917, 265020, 265021, 52644150, 264693, 35695855, 264632, 52644332, 22279002
2406		111, 4812	88094501 (4811, 4812) Novel Protein sim. GBank gi 2773363 (AF041382) - microtubule binding protein D-CLIP-190 (Drosophila melanogaster]	CAP-Gly domain CAP-Gly domain	struct	52646842, 22278994, 22278996, 35696286, 22278997, 22278999, 22478999, 284092, 264093, 60432049, 264259, 29331822, 29331827, 3569652, 29331828, 29331827, 3569652, 29331828, 264106, 264908, 265010, 265007, 265008, 265009, 264310, 265010, 265011, 265018, 265019, 264319, 2648150, 2648150, 2648139, 264823, 264628, 35698423, 264628, 35698423, 264639, 22278000, 22278000, 26278000, 26278000, 26278000, 26278000, 26278000, 26278000, 26278000, 26278000, 26278000, 26278000, 264563
2407		313, 4814			UNCLASSIFIED	264685, 264686
2408		315, 4816	87391503 (4815, 4816) Novel Protein sim. GBank gil423442 pir  S33513 - gene Fif protein - mouse		UNCLASSIFIED	264910, 265010, 264448, 264557

22278995, 22278996, 22278997, 264097, 29331822, 29331827, 29146498, 52644045, 60433438, 33657084, 87168474, 264760, 21906767, 29148627, 29148629, 52644150, 33657023, 263967, 20281069, 18108374, 20281071, 56182323, 83373044, 18108385, 87168518	22278998, 264259	29331824, 29331827, 29331828, 264764, 264369, 33657109, 56182323	22278998, 29331827, 264907, 265011, 265017, 265018, 265019, 18108351, 21906768, 265020, 33657109, 264559, 18108385	264488, 264259, 29331826, 264508, 264905, 264509, 264906, 264507, 264908, 264906, 264510, 264511, 264512, 265008, 265009, 264910, 265011, 264682, 264764, 264768, 264686, 264768, 264689, 266685, 2646370, 264637, 264636, 35695855, 264632, 264564, 264565, 264566, 264567, 264486	52645156, 52646842, 52646305, 18108398, 526182575, 22278994, 22278995, 56994075, 22278996, 326986075, 22278996, 326986075, 22278998, 326986075, 22278999, 22278998, 32694075, 22278999, 22278999, 22278999, 22278999, 22331822, 29331824, 29331827, 29331828, 2564507, 29331827, 246699, 60433356, 3365790, 284590, 264684, 264687, 264688, 56181562, 21906764, 264689, 21906765, 21906765, 21906767, 29148627, 21906765, 21906766, 21906767, 29148627, 21906769, 264693, 265021, 60170615, 33657039, 264693, 25645129, 33657109, 33657182, 27486261, 27486265, 33657349, 18108376, 60431528, 264629, 18108374, 18108336, 284558, 5618223, 83373044, 18108335, 22278002, 2265180, 22278002, 2265180, 22278002, 2265180, 22278002, 22278002, 2265180, 22278002, 22272002, 222720, 222
UNCLASSIFIED		tnf	dna_rna_bind		ubiquilin
<b>.t.</b> .	Contains protein domain (PF00080) - Copper/zinc superoxide dismutase (SODC)		Contains protein domain (PF00076) - Idna_rna_bind RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)	Contains protein domain (PF00038) - struct Intermediate filament proteins	
Novel Protein sim. GBank gil 176601 sp P45966 YNZ6_CAEEL - HYPOTHETICAL 20.8 KD PROTEIN T09A5.6 IN CHROMOSOME III	87604860 (4819, 4820) Novel Protein sim. GBank gil4966262lgb AAC48052.2  - (U64849) Contains similarity to Pfam domain: PF00646 (F-box), Score=28.7, E-value=4.3e-05, N=1 [Caenorhabdilis elegans]	87534633 (4821, 4822) Novel Protein sim. GBank gij3114713 (AF061346) - Edp1 protein [Mus musculus]	87778332 (4823, 4824) Novel Protein sim. GBank gil5410336 gb AAD43038.1 - (AF106685) myelin gene expression factor 2 [Homo sapiens]	94133820 (4825, 4826) Novel Protein sim. GBank gil5262705 emb CAB45778.1  - (AL080214) hypothetical protein [Homo saplens]	94312590 (4827, 4828) Novel Protein sim. GBank gij1082340[piri  S52863 - DNA-binding protein R kappa B - human
2409   94741770 (4817, 4818)   Novel Protein sim. GBank gil1176601lsp P45966 YN 20.8 KD PROTEIN T09A5	87604860 (4819, 4820)	_		94133820 (4825, 4826)	94312590 (4827, 4828)
2409	2410	2411	2412	2413	2414

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264259, 264908, 60433356, 33657402, 21906754, 265018, 264687, 264689, 21906769, 55811957, 265021, 264690, 264691, 33657023, 264693, 35696423,	264638	264094, 29331822, 29331824, 29331827, 264369	56994075, 264091, 264259, 29331824, 29331825, 60432289, 29331828, 264905, 29331828, 264905, 264907, 264907, 265009, 60432229, 21906754, 87168559, 265019, 264682, 21906769, 265020, 265021, 33657023, 65274620, 18108385, 22279000, 22279002, 264534, 18108385, 22279000, 22279002, 264566	264569, 264762, 264448, 264691, 264631, 264634, 264555, 264556, 264638, 264558	29331824, 29331825, 29331828, 60432229, 33109954, 85658542, 87168474, 265018, 264288, 2650 <u>20, 28456</u> 4	264909, 264768, 264638	18108392, 18108394, 18108399, 269905, 265006, 265010, 18108351, 18108374, 18108385	264259, 60432049, 29331822, 29331826, 60432289, 29331828, 265008, 265009, 6043336, 21906754, 265017, 265018, 265019, 21906766, 21906768, 21906769, 265020, 265021, 20281149, 263971, 60432113	5618257, 33089460, 74270357, 0541827, 2641250, 26331826, 29331828, 264905, 265110, 26519, 264766, 21906768, 55811957, 264692, 33657023, 33657109, 55811576, 56182323, 83373044, 18108385, 18108388, 60432113, 22279000	264634	22278996, 22278998, 264259, 264102, 264512, 265008, 21906767, 18108370, 18108374, 263976
			UNCLASSIFIED	struct	UNCLASSIFIED	UNCLASSIFIED	transport	kinase	UNCLASSIFIED	kinase	UNCLASSIFIED
423915 pir  A45439 - myosin   Contains protein domain (PF00063) - Struct Myosin head (motor domain)		Contains protein domain (PF00153) - Mitochondrial carrier proteins	Contains protein domain (PF00411) - UNCLASSIFIED Ribosomal protein S11				Contains protein domain (PF00617) - transport RasGEF domain	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain		Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	
2415   88088002 (4829, 4830) Novel Protein sim. GBank gil423915 pir  A45439 - myosin l heavy chain - rat	94118356 (4831, 4832) Novel Protein slm. GBank gij3025445 (AC004528) -	R32184 1 Homo sapiens/ 8773334 (4833, 4834) Novel Protein sim. GBank gi1084944[pirl]S54495 - hypotelical protein YPR021c - yeast (Saccharomyces	Catewards) Novel Protein sim. GBank gij1176572 sp P45895 YNA4_CAEEL - HYPOTHETICAL 91.0 KD PROTEIN PAR2.4 IN CHROMOSOME III	82374249 (4837, 4838) Novel Protein sim. GBank gi 284006 pir  518732 - aufoantigen, 64K - human	94844244 (4839, 4840) Novel Protein sim. GBank gil1076211 pir  S50755 - hypothetical protein VSP-3 - Chlamydomonas reinhardtii		88084714 (4843, 4844) Novel Protein sim. GBank gi[2224567]dbj BAA20772  - (AB002311) KIAA0313 [Homo sapiens]		2424   94854047 (4847, 4848) Novel Protein sim. GBank gi[2988398 (AC004381) - Unknown gene product [Homo sapiens]	87415981 (4849, 4850) Novel Protein sim. GBank gip2077932 dbj BAA19879  - IOR8556) Protein Kinase Rattus noveolous	
88089002 (4829, 4830) N	94118356 (4831, 4832) N	87733334 (4833, 4834) N	94234349 (4835, 4836) Novel Protein sim. GBank gil116572 sp P45895 YN 91.0 KD PROTEIN PAR2.	82374249 (4837, 4838) h	94844244 (4839, 4840)	87805345 (4841, 4842)	88084714 (4843, 4844)	88058390 (4845, 4846)	94854047 (4847, 4848)		
2415	2416	2417	2418	2419	2420	2421	2422	2423	2424	2425	2426

•	•										
264259, 20281099, 35696052, 265008, 264584, 265011, 264760, 18108351, 264682, 264683, 264684, 264686, 264687, 264689, 21908766, 264691, 264692, 18108374, 18108377, 264557, 264639,	22278996, 22278999, 35696052, 21906754, 264288, 21906765, 21906768, 21906769, 35695917, 265020, 263972, 22279002	22278999, 29331824, 264906, 264909, 264511, 265009, 21906754, 265017, 265018, 265019, 264448, 264683, 264288, 21906765, 21906768, 265021, 264693, 18108381	264112, 264691	29331826, 29331827, 35696052, 29146499, 264905, 264906, 264681, 264288, 264689, 21906765, 264692, 35696423	264634, 264558	264569, 264905, 265018, 264762, 264683, 264591, 264556, 264557, 264639, 264558	264563	264555	265008, 264758, 265010, 264689, 27486261, 263972, 18108374, 18108381	29331828, 265007, 265009, 265017, 264760, 264685, 264693, 264565	29331826, 264508, 264905, 264509, 264906, 264907, 264908, 264906, 264511, 265006, 264512, 264910, 265009, 264511, 265006, 264512, 264910, 265009, 264764, 265011, 264760, 264764, 264585, 264693, 264591, 30557023, 264693, 264531, 264532, 564565, 264565, 264565, 264567
ribosomalprot		ÜNCLASSIFIED	UNCLASSIFIED		protease	transcriptfactor		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED
Contains protein domain (PF00573) - inbosomalprot Ribosomal protein L4/L1 family		Contains protein domain (PF00711) - UNCLASSIFIED Beta defensins		Contains protein domain (PF01728) - FtsJ cell division protein		Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type					Contains protein domain (PF01256) - UNCLASSIFIED Uncharacterized protein family UPF0031
2427   87622693 (4853, 4854) Novel Protein sim. GBank gij4680695[gb]AAD27737.1 AF13296 - (AF132962) CGI-28   F protein [Homo sapiens]	85732889 (4855, 4856) Novel Protein sim. GBank gi[1537070 (U63840) - Inucleoporin p54 [Rattus norvegicus]	Novel Protein sim. GBank gi[601931 (M94316) - neurofilament-H [Oryctolagus cuniculus]	3, 4860)	Novel Protein sim. GBank gij3860729jemb CAA14630j - (AJ235270j CELL DIVISION PROTEIN FTSJ (flsJ) [Rickettsia prowazekii]	80083033 (4863, 4864) Novel Protein sim. GBank gij3976367 emb CAA93287  - (Z69360) Weak similarity to Eimeria thrombospondin (PIR Acc. No. A45517); cDNA EST EMBL:M89266 comes from this gene; cDNA EST yk295b9.5 comes from this gene   [Caenorhabditis elegans]	k gi[2224593]dbj[BAA20784] - tomo sapiens]		9, 4870)	88044008 (4871, 4872) Novel Protein sim. GBank gi 1263289 (U47856) - fibroin-4 [Araneus diadematus]	83363424 (4873, 4874) Novel Protein sim. GBank gij3641352 (AF091234) - putative Iranscription factor (Mus musculus)	94143473 (4875, 4876) Novel Protein sim. GBank gi 3860014 (AF091088) - unknown [Homo sapiens]
427 87622693 (4853,	2428 85732889 (4855,	2429 87769276 (4857,	2430 86948827 (4859, 4860)		2432   80083033 (4863,	2433 80055092 (4865,	$\Box$	2435  20759044 (4869, 4870)	2436 88044008 (4871,	2437   83363424 (4873,	2438   94143473 (4875,

60424179, 18108397, 56182575, 56181686, 56994075, 22278996, 35696286, 22278997, 22278999, 2642596, 25645080, 29331822, 56182181, 29331824, 60424269, 66714117, 29331825, 60432289, 29331826, 29331827, 29331828, 264308, 264308, 264308, 264308, 264308, 264308, 264308, 264308, 264308, 26431735, 33109954, 21906754, 265017, 265018, 265019, 18108351, 265017, 265018, 265019, 18108351, 264448, 264288, 264768, 2564429, 56181562, 21906764, 21906765, 21906766, 21906766, 21906766, 21906766, 21906766, 21906766, 21906766, 21906767, 265018, 2569556, 18108370, 60431528, 263977, 55810764, 35696423, 6643113, 22279000, 22279002, 264567	29331822, 29331824, 29331825, 29331827, 35696052, 264508, 264907, 264510, 265018, 265019, 265019, 265023, 265023, 264639, 22279002	264488, 264629, 18108374, 264564	22278996, 22278999, 29331822, 264768, 264693	264906	264259, 18108382, 18108383, 18108385, 22279000	264259, 35696052, 264369, 18108361	265011, 264689, 33657023, 263981, 18108385	56182575, 264259, 29331824, 29331825, 29331827, 60433356, 60433438, 264758, 265018, 264692, 65274620, 60431528, 65274791, 56182323
UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	Iranscriptfactor	transcriptfactor		proteaseinhib	cadherin
Contains protein domain (PF01423) - UNCLASSIFIED Sm protein			Contains protein domain (PF00560) - UNCLASSIFIED Leucine Rich Repeat	Contains protein domain (PF01352) - KRAB box	Contains protein domain (PF00170) - Iranscriptfactor bZIP transcription factor		Contains protein domain (PF00095) - proteaseinhib WAP-type (Whey Acidic Protein)  Your-disulfide core*	
Novel Protein sim. GBank gil4263519[gb AAD15345] - (AC004044) small nuckear riboprotein Sm-D1 [Arabidopsis thaliana]	1880)	87623914 (4881, 4882) Novel Protein sim. GBank gij3024889 sp P56524 Y288_HUMAN - HYPOTHETICAL PROTEIN KIAA0288 (HA6116)	87273590 (4883, 4884) Novel Protein sim. GBank gil4505013[ref]NP_002703.1[pPPP1 - protein phosphalase 1. regulatory subunit 7	84305949 (4885, 4886) Novel Protein sim. GBank gi 1170658 sp Q02975 KID1_RAT Contains protein domain (PF01352) - Iranscriptfactor - RENAL TRANSCRIPTION FACTOR KID-1 (TRANSCRIPTION FACTOR 17)	88086345 (4887, 4888) Novel Protein sim. GBank gil4758824[ref]NP_004280.1[pNRF3 - nuclear factor [erythrold-derived 2]-like 3	87338636 (4889, 4890) Novel Protein sim. GBank gij2135950 pir  S58222 - PQ-rich   protein - human	88059293 (4891, 4892) Novel Protein sim. GBank gil4753887 emb CAA05409.2  - (AJ002424) p65 protein [Rattus norvegicus]	94845149 (4893, 4894) Novel Protein sim. GBank gil4885613 ref NP_005409.1 pST5  - suppression of tumorigenicity 5
99 94850850 (4877, 48			2442   87273590 (4883, 48	2443 84305949 (4885, 48				2447   94845149 (4893, 48

2448	2448 87749680 (4895, 4896)			UNCLASSIFIED	22278996, 22278997, 22278999, 29331826, 35696052, 264107, 264110, 87168474, 87168559, 18108351, 21906767, 21905769
					27486262, 263976
2449	87869075 (4897, 4898)	87869075 (4897, 4898) Novel Protein sim. GBank gij7288371spP39194/ALU7_HUMAN - IIII ALU SUBFAMILY		cadherin	264259, 264828, 265007, 264595, 265021, 56526486
2460	10000 4000	OC WARMING ENIKT III			300736
2664	00031 104 (4683, 4800)	The state of the s	11 COCOTO, -1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1	UNCLASSIFIED	204900
643	91014503 (4901, 4902)	91014303 (4901, 4902) Novel Protein Sim. GBank gi 1710021 sp P35290 RB24_MOUSE - RAS-RELATED PROTEIN RAB-24 (RAB-16)	Contains protein domain (Prudu / 1) - glycoprotein Ras family	glycoprotein	264093, 29331822, 29331824, 29331825, 66714117, 29331826, 29331828, 35696052, 264907, 66712502, 29331830, 264910, 265009, 264758, 265017, 265018, 264762,
					264448, 264288, 21906767, 265021, 33657023, 264693, 33657109, 263969,
					83373044, 18108385
2452	91230509 (4903, 4904)	91230509 (4903, 4904) Novel Protein sim. GBank gi 1504034 db  BAA13216  -  (D86980) KIAA0227 [Homo sapiens]		isomerase	264102, 264112, 264688, 263972, 18108374, 83373044, 264563
2453	84201088 (4905, 4906)	84201088 (4905, 4906) Novel Protein sim. GBank gij2880079 (AC004142) - similar	Contains protein domain (PF00560) - ngfrecep	ngfrecep	264509, 264512, 18108385
		to murine leucine-rich repeat protein; possible role in neural development by protein-protein interactions; 93% similarity to D49802 (PID:91369906) [Homo sapiens]	Leucine Rich Repeat		
2454		95310691 (4907, 4908) Novel Protein sim. GBank gil1076802 pir  S49915 - extensin	gi 1076802 pir  S49915 - extensin Contains protein domain (PF00170) -  UNCLASSIFIED	UNCLASSIFIED	263994, 66714117, 29331827, 264508,
		like protein - maize	bZIP transcription factor		264509, 264905, 264906, 264907, 264908.
					264909, 264510, 264511, 264512, 265009.
		-			264910, 264591, 264758, 264759, 265010,
					265011, 264603, 264604, 264760, 264761,
					264762, 18108351, 264764, 264765, 264766,
					264686, 264768, 264769, 264534, 264691,
					264692, 33657023, 264693, 33857109,
					264628, 263978, 35695855, 264634, 264635,
			,		264637, 264638, 264639, 83373044,
	_				18108385, 264563, 264564, 264486
2455	-	95288301 (4909, 4910) Novel Protein sim. GBank	•	glycoprotein	264488, 22278996, 264259, 35696052,
		gij543817ispiP35585jAP47_MOUSE - CLATHRIN COAT	Adaptor complexes medium subunit		264905, 264906, 264907, 264908, 264909,
		ASSEMBLY PROTEIN AP47 (CLATHRIN COAT	family		264510, 264512, 265008, 265009, 264910,
	-	ASSOCIATED FACTOR SITUATION ACTION AND SITUATION AT THE STATE OF THE SITUATION AND SIT			204391, 204332, 204333, 204334, 204339, 366010 364760 364881 18108351 364683
		ASSEMBLY PROTEIN ASSEMBLY PROTEIN COMPLEX 1			200013, 204100, 204001; 10100031; 204003; 1084764 18108354 264766 264768 264769
		MEDIUM CHAIN			264689, 21906766, 21906767, 21906769,
					29148629, 35695917, 265020, 265022,
	-				33657023, 33657109, 18108370, 264628,
					264629, 264631, 264632, 264635, 56182323,
			_		60170394, 18108385, 264563, 264564,
	_				264566, 264567
2426		88166700 (4911, 4912)  Novel Protein sim. GBank gil2588630 (AC003079) - Ankyrin like; 54% similar to 2022340A (NID:g1092123) in exons spaning 43074 to 14551 of plant Home	gil2588630 (AC003079) - Ankyrin Contains protein domain (PF00023) - kinase 40A (NID:g1092123) in exons Ank repeat	kinase	264693
	1	spanning 43974 to 11331 of done, [homo saprens]			

2457	94118375 (4913, 4914)	2457   94118375 (4913, 4914) Novel Protein sim. GBank gij3025447 (AC004528) -   R32184_3 [Homo sapiens]		UNCLASSIFIED	56181686, 264905, 264907, 264511, 264596, 55811386, 264687
					264691, 33657023, 264693, 35695855,
					264636, 264555, 56182323, 264558. 56526486, 264563
2458		85675304 (4915, 4916) Novel Protein sim. GBank gil2384942 (AF022985) - Similar Ito collagen [Caenorhabditis elegans]		UNCLASSIFIED	264691, 264693, 264634, 264559
2459			Contains protein domain (PF00097) - transport	transport	27486265
		gips4 1942ggpAAD43187.1AC00489 - (AC004897) supported by mouse EST AA538043 (NID:g2284036) Homo caniens	Zinc finger, C3HC4 type (RING finger)		
2460	94315289 (4919, 4920)	94315289 (4919, 4920) Novel Profein sim GRank		900013	
		gi4929701fablAAD34111,1/AE15187 - (AE151874) CGI-116		RINASG	032/43/4, 33696286, 222/8996, 222/899/, 60433046, 30334832, 30334834, 30334836
		protein [Homo sapiens]			66714117, 60432289, 29331826, 29331827,
					29331828, 33656970, 29146499, 264102,
					264109, 60433438, 265017, 265018, 265019,
				-	264288, 21906765, 21906766, 21906769,
					35695917, 265020, 264691, 33657023,
					27486261, 18108374, 35695855, 87168518.
2481	87646147 (4021 4022)				60432113
	(4821, 4822)	0/04314/ (4861, 4862) Novel Protein sim. GBank gij4426962 gb AAD20633  -		UNCLASSIFIED	264259, 29331828, 264910, 18108351,
		sapiens]			18108370, 18108374
2462	86998002 (4923, 4924)	86998002 (4923, 4924) Novel Protein sim. GBank gij5420387 emb[CAB46679.1] -			264909 264758 264684 18108374 264637
_[		(AJ243459) proteophosphoglycan (Leishmanta major)			18108385
7403	84388543 (4925, 4926)		Contains protein domain (PF00096) - UNCLASSIFIED	Γ	264681, 264566
			Zinc finger, C2H2 type		
2464	91219957 (4827, 4928)	D43021.11-	Contains protein domain (PF01399) - protease	protease	264489, 52646842, 22278995, 35696286
			PCI domain		22278996, 22278997, 22278999, 264259,
	-				29331822, 29331824, 66714117, 29331825.
					29331826, 29331827, 35696052, 29331828,
					264509, 52644045, 264510, 264511, 264512.
					265008, 60170831, 264593, 52646317,
					33109954, 33657084, 265017, 265018,
					265019, 264762, 264448, 264764, 264288,
					264766, 21906765, 21906766, 21906767.
					21906768, 21906769, 265021, 33657023,
					33657109, 18108370, 18108381, 60170394,
					18108385, 22279002, 264486

2465	953574B3 (4929, 4930)	2465   95357483 (4929, 4930) Novel Protein slm. GBank gi 4508401 ref NP_002871.1 pRAF1 - v-raf-1 murine leukemia viral oncogene homolog 1	Contains protein domain (PF00069) - oncogene Eukaryotic protein kinase domain		18108392, 52644507, 52645156, 52646365, 22278994, 22278995, 35696286, 22278996, 22278999, 22278995, 35696286, 22278996, 22278998, 264259, 29331822, 29331824, 29331825, 6442229, 29331824, 35696052, 29331828, 264907, 29331830, 526907, 255008, 265009, 264910, 33657402, 265010, 265010, 265011, 87168559, 264910, 33657402, 265010, 265011, 87168559, 264600, 265017, 265018, 265019, 18108351, 264369, 264288, 264685, 264767, 21906765, 21906767, 21906768, 55611957, 35695817, 265020, 265021, 60170615, 52644150, 33657023, 35695763, 264628, 264637, 61170394, 35695464, 264566, 264887
		85681386 (4931, 4932) Novel Protein sim. GBank gil4321619 gb AAD15788.1  - (AF051098) seven transmembrane domain orphan receptor [Mus musculus]			264369
2467		88059465 (4933, 4934) Novel Protein sim. GBank gi]3513300 (AC005595) - F16601_1, partial CDS [Homo sapiens]		UNCLASSIFIED	56994075, 264908, 21906768, 33657023
2468		Novel Protein sim. GBank gi 2143455 pir  I58106 - gene DMR-N9 protein - mouse (fragment)	Contains protein domain (PF00400) - kinase WD domain, G-beta repeat		29331824, 52644045, 265008, 264910. 265019, 21906765, 21906769, 265021
2469	_				264288, 264628
2470		80223831 (4939, 4940) Novel Protein sim. GBank gij5420389jembjCAB46680.1j - (AJ243460) proteophosphoglycan (Leishmania major)		UNCLASSIFIED	264592, 264692, 264555, 264556, 264557, 264558, 264559, 18108385, 264482
2471		91013681 (4941, 4942) Novel Protein sim. GBank gi 5419882 emb CAB46424.1  - (AL096749) DKFZp434G153 [Homo sapiens]		UNCLASSIFIED	65274572, 35696286, 29331827, 265007, 264592, 33109954, 265018, 265019, 264288, 55811957, 265020, 264693, 55811576, 56182323
2472		95050811 (4943, 4944) Novel Protein sim. GBank gil4929747[gb]AAD34134.1 AF15189 - (AF151897) CGI-139 protein [Homo sapiens]		UNCLASSIFIED	264092, 60432049, 29331825, 60433356, 265010, 265011, 18108351, 264784, 264288, 264692, 65274620, 18108370, 18108372, 18108374, 264634, 18108385
2473		95421509 (4945, 4946) Novel Protein sim. GBank gil4539009jembjCAB39630.1j - (AL049481) putative protein [Arabidopsis thaliana]			60424179, 65274572, 22278999, 60424269, 29331826, 265008, 60433356, 60433438, 265010, 18108351, 284448, 264288, 264687, 264689, 265021, 264692, 65274620, 60431528, 65274791, 264556, 56182323, 60432113
2474		94315616 (4947, 4948) Novel Protein sim. GBank gi]3252827 (AC004382) - Unknown gene product [Homo sapiens]		***	65274572, 56994075, 264259, 29331826, 60170831, 265017, 265018, 265019, 264683, 264369, 265020, 264693, 264563, 264564

2478   9421653 (4949, 4950)   Novel Protein sim, GBank gilt216466 (U48852) - HT protein Contains protein domain (FF00009) - 1g1	264259, 29331822, 265006, 265007, 265010,	265011, 264448, 264288, 264389, 264885, 264686, 18108357, 264768, 18108362	264693, 18108370, 18108374, 18108379.	35696423, 83373044, 18108383, 18108385,	264564, 264565, 264567	264259, 60424269, 66714117, 264905,	265006, 264511, 265008, 265009, 264758.	265010, 265011, 18108351, 264681, 264369,	284288, 264689, 21906767, 265020.	18108374, 264639, 18108382, 83373044,	18108385 87158518	28.307.8	0,6207	76501/	56994075, 22278999, 21906754, 264682. 21906765	264905, 264907, 264765	65274572, 56182575, 22278997, 264094.	264259, 29331822, 29331824, 66714117.	29331827, 35696052, 264508, 264905,	264908, 264907, 264908, 52644045, 264909,	58182435, 265008, 264910, 33657402.	55812038, 264758, 265010, 265011, 265017,	265018, 264760, 264762, 18108351, 264764,	264288, 264766, 264686, 284768, 21906768,	55811957, 265020, 264691, 264692, 264693,	264629, 55811576, 264630, 264634, 264635,	264636 264637 264556 264558 56182323.	83373044, 60432113, 22279002	29331822, 29331824, 29331825, 29331827,	264508 264905 264509 264906 264907.	264908, 264511, 264591, 264768, 264693,	264631, 264632, 264636, 264638, 264639.	264563	264488, 22278995, 264093, 264095,	60432049, 60433356, 60433438, 264448.	264288, 263967, 18108370, 18108385,	18108388, 264482	264563	22278995, 22278996, 22278997, 22278999,	264259, 60432049, 29331824, 60432289.	29331827, 265007, 264910, 264593, 264600.	264603, 264604, 265019, 264448, 264288.	264685, 264686, 264769, 264689, 35695917.	265022, 264692, 264693, 56182323
121693 (4949, 4950)   Novel Protein sim. GBank gill 216496 (U49552) - HT protein Contains protein domain (PF00009)   Criceulus griseus    Criceulus griseu	lgí					UNCLASSIFIED						I INC. ACCIEIED	UNCLASSIFIED	UNCLASSIFIED	tm7	UNCLASSIFIED	collagen				•	•							UNCLASSIFIED					UNCLASSIFIED				kinase	UNCLASSIFIED					
Cricetulus griseus	Contains protein domain (PF00008) -	EGF-like domain																																										
315618 (4951, 4952) 718974 (4953, 4954) 559165 (4955, 4956) 314569 (4957, 4958) 718481 (4961, 4962) 718481 (4961, 4962) 731583 (4965, 4966) 731583 (4965, 4966)	Novel Protein sim. GBank gil 1216486 (U48852) - HT protein	[Cricetulus griseus]				Novel Protein sim. GBank gij3252827 (AC004382) -	Unknown gene product [Homo sapiens]								Novel Protein sim. GBank gi 1644232 db  BAA11082  -  D67066  N-WASP IBos taurus		Novel Protein sim. GBank gil5689469IdbjlBAA83018.11 -	(AB028989) KIAA1066 protein [Homo sapiens]		•									Novel Protein sim GBank nil321249lnirllS28407 - quanine	Inicipolitide exchange activator CDC25 homolog - mouse	BODON - ROMANIA CARONO CORANAS ORIGINAS - CONCARAS.								Novel Protein sim. GBank gil1185397 (U25281) - SH3	domain binding protein (Rattus norvegicus)				
2475     945       2476     946       2477     20       2477     20       2481     94       2482     87       2483     87       2484     94       2485     87	94321693 (4949, 4950)					94315618 (4951, 4952)						_	_	1	94314569 (4957, 4958)	95295605 (4959, 4960)	_																	87731583 (4985, 4966)			~							

265017, 264555	22278996, 35696286, 22278997, 22278998, 22278999, 264092, 264259, 28331822, 35696052, 264108, 264269, 28331822, 35696052, 264108, 264905, 264907, 264907, 265008, 26433438, 21906754, 33109954, 87188559, 265018, 264681, 264288, 264687, 21906785, 21906768, 21906768, 21906768, 21906769, 21906769, 35695917, 265021, 26502, 264534, 33657023, 264692, 33657109, 263972, 18108377, 3569643, 33657109, 22279002, 264563, 264482, 264565, 202279000, 22279002, 264563, 264482, 264565,	UNCLASSIFIED 264448, 264288, 264684, 264691, 264631,	nplementrecept 264688, 264693, 55811576, 22279002	UNCLASSIFIED 264907, 265008, 22279002	7 264259, 29331826, 265008, 264762. 18108370, 18108376, 18108379	transcriptfactor 264488, 22278996, 22278999, 29331828, 264591, 33109954, 265017, 55811150, 21906768, 21906768, 264692, 60431528,
	gij464559jspjP35287jRB14_RAT Contains protein domain (PF00071) - glycoprotein Ras family	5	Contains protein domain (PF00084) - complementrecept Sushi domain (SCR repeat)		CU1	, , , , , , , , , , , , , , , , , , ,
Novel Protein sim. GBank gi 2662167 dbj BAA23715  - AB007903) KIAA0443 [Homo sapiens]	95343105 (4973, 4974) Novel Protein sim. GBank gil464559 sp P35287 RB14_RAT RAS-RELATED PROTEIN RAB-14		82990585 (4977, 4978) Novel Protein sim. GBank gil4886439[emb CAB43355.1] - [AL050253) hypothetical protein [Homo saplens]	2490   88069609 (4979, 4980) Novel Protein sim. GBank gi(2588624 (AC003083) - Rap2 interacting protein-like; similar to U73941 (PID:g1916018) [Homo sapiens]	91242116 (4981, 4982) Novel Protein sim. GBank gil728832[sp P39189[ALU2_HUMAN - III! ALU SUBFAMILY SB WARNING ENTRY IIII	95308202 (4983, 4984) Novel Protein sim. GBank gil3355303 (AF001549) - Unknown gene product [Homo sapiens]
2486   87748978 (4971, 4972)   Novel Protein slm. GBank (   (AB007903) KIAA0443   Ho	95343105 (4973, 4974) R	87652451 (4975, 4976)	82990585 (4977, 4978) I	88069609 (4979, 4980)	91242116 (4981, 4982)	95308202 (4983, 4984)
2486	2487	2488	2489	2490	2491	2492

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sinct	I INCI ASSIGIED	UNCLASSIFIED	Collage				struct		UNCLASSIFIED			UNCLASSIFIED	9) - kinase
Contains protein domain (PF01424) - Struct					•					1			Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain
2493   95422415 (4985, 4986) Novel Protein sim. GBank gil4240307 dbj BAA74932.1  - (AB020716) KIAA0909 protein [Homo sapiens]		Marri Descript of the Court of 642003001	94234331 (4969, 4990) Nover Protein sim. GBank gij342039jembjCAB46680.1] - (AJZ43460) proteophosphoglycan [Leishmania major]				80018765 (4991, 4992) Novel Protein sim. GBank gil4808220jembjCAB42832.1j -	(AL022315) dJ117715.1 (PUTATIVE novel protein) [Homo sapiens]				87724633 (4995, 4996) Novel Protein sim. GBank gil 1200503 (U47924) - B [Homo sapiens]	94685125 (4997, 4998) Novel Protein sim. GBank gi[3510234 (AC005581) - R31237_1, partial CDS [Homo sapiens]
3 95422415 (4985, 4986)	34 30703118 (4087 4088)	2406 04224664 (4080 4000)	32 84234551 (4868, 4880)				2496 (80018765 (4991, 4992)		2487 91723554 (4993, 4994)			2488 87724633 (4995, 4996)	2499   94685125 (4997, 4998)

2504 87868708 (5007, 5008) Novel Protein sim. GBank gil550420jembjCAA48220j -			264488, 52644507, 52645156, 52646842. 22278994, 264259, 52645080, 29331822,
Sprifiano			29331824, 29331825, 29331826, 29331827, 35696052, 264906, 264908, 52644045, 265009, 60433356, 33657402, 60433438, 264595, 33109954, 87168474, 265017, 265019, 264448, 264288, 264766, 52644229,
			21906765, 21906766, 21906767, 21906768. 52644150, 264692, 27486261, 27486262, 27486264, 27486265, 35695763, 35696423,
		UNCLASSIFIED	3583583, 32843352, 30182523, 10182533, 1018253, 1018253, 1018253, 1018253, 1018253, 1018253, 10182533, 1018253, 1018253, 1018253, 1018253, 1018253, 1018253, 10182533, 1018253, 1018253, 1018253, 1018253, 1018253, 1018253, 10182533, 10182533, 10182533, 10182533, 10182533, 10182533, 10182533, 10182533, 10182533, 10182533, 10182533, 10182533, 10182533, 101
8760559 (3009, 3019) 91232328 (5011, 5012) Novel Protein sim. GBank gi[2137562]pirj[149635 - mouse	und		264488, 52644507, 52645156, 52646365, . 65274572, 22278995, 56994075, 22278996, (
Dhm1 protein - mouse			22778997, 22278999, 22278999, 264259, 60432049, 29331822, 29331825, 29331825, 29331826, 29331828, 264509, 56182435, 264112, 264593, 60433358, 55812038, 21906754, 265011, 265017, 265018, 265019, 264605,
			264762, 18108351, 26448, 264288, 264768, 21906765, 21906766, 21906767, 21906768, 21906769, 35692917, 265020, 265022, 60170615, 33657023, 27486264, 18108379, 35695855, 264637, 83373044, 18108365, 87168518, 60432113, 22279000, 264563, 264585
95316233 (5013, 5014) Novel Protein sim. GBank gil5174489[ref]NP_006035.1 pKIAA - histone deacetylase 6 Hi	Contains protein domain (PF00850) - histone Histone deacetylase family	stone	264488, 263994, 284592, 264595, 264369, 264686, 264768, 35695917, 35696423, 264563
95315505 (5015, 5016) Novel Protein sim. GBank gil4826433 emb CAB42889.1  - (AL031447) dJ126A5.2.1 (novel protein) (isoform 1) [Homo sapiens]	5	UNCLASSIFIED	22278995, 22278999, 60432049, 284259, 29331828, 285006, 265007, 60433438, 33657084, 265010, 265017, 265018, 265019, 18108351, 264448, 18108354, 284369, 18108359, 21906765, 21906769, 55811957, 265020, 265022, 27486261, 33657349, 18108377, 35695855, 60432113, 22279002, 264563, 264565
87813741 (5017, 5018) Novel Protein sim. GBank gij1263289 (U47856) - fibroin-4  Araneus diadematus	5	UNCLASSIFIED	265007, 265008, 18108357, 264556, 264567

65274572, 22278994, 22278996, 22278998, 22278999, 60432049, 264259, 29331822, 29331824, 25331825, 66714117, 60432289, 22331828, 33558970, 264908, 66712502, 265007, 264910, 60170831, 60433229, 60433356, 6043348, 21906754, 87168474, 265017, 265018, 26448, 264288, 21906767, 21906768, 21906769, 25448, 265020, 265022, 264691, 33657023, 264693, 65274620, 33657109, 33657182, 27486262, 33657349, 18108370, 3569565, 264555, 26182323, 83373044, 60432113, 22279002	265017, 21906764, 265020, 264692	22278999, 264259, 29331825, 29331826, 29146499, 264907, 264909, 265006, 265008, 264591, 60432229, 21906754, 264763, 264683, 264766, 18108357, 264689, 264693, 18108370, 263972, 18108374, 264558, 22279000	60424179, 52645156, 18108394, 22278994, 35696286, 56994075, 22278996, 29331822, 29331824, 60424269, 29331825, 29331827, 33656970, 60431735, 33657084, 87168559, 265017, 264448, 264369, 56181562, 21906766, 21906766, 21906769, 21906768, 21906769, 265020, 265031, 33657023, 18108386, 18108374, 55810764, 35696423, 56182323, 264558, 18108385	264510		264259, 29146498, 264905, 264288, 29148629, 35695917, 27486261, 264634	264091, 29331824, 29331825, 29331826, 29331828, 29331828, 35696052, 264508, 264509, 264905, 264909, 264511, 264910, 33657402, 264757, 33109954, 265017, 265018, 264766, 264769, 33657109, 33657109, 3565782, 264768, 264768, 264639, 35657182, 264628, 55811576, 35696423, 264631, 264634, 264637, 264638, 264564
dna_ma_bind		transport	UNCLASSIFIED	UNCLASSIFIED	tm7	UNCLASSIFIED	UNCLASSIFIED
		Contains protein domain (PF00097) - transport Zinc finger, C3HC4 type (RING finger)			Contains protein domain (PF00001) - Im7 7 transmembrane receptor (rhodopsin family)		
Novel Protein sim. GBank gij3293537 gb AAC25762.1  - (AF071059) zinc finger RNA binding protein [Mus musculus]	87384281 (5021, 5022) Novel Protein sim. GBank gil4323152 gb AAD16228.1  -   (AF098863) Ets-protein Spi-C [Mus musculus]	88084771 (5023, 5024) Novel Protein sim. GBank gil4502075 refiNP_001135.1 pAMFR - autocrine motility factor receptor	95357843 (5025, 5026) Novel Protein sim. GBank gij3004657 (AF017777) - bobby sox [Drosophila melanogaster] '	88094578 (5027, 5028) Novel Protein sim. GBank gij2258437 (AF008197) - syncollin (Rattus norvegicus)			87784966 (5033, 5034) Novel Protein sim. GBank gil4220527 emb CAA23000  - (AL035356) putative protein [Arabidopsis thaliana]
2510   95421379 (5019, 5020)   Novel Protein sim. GBank g (AF071059) zinc finger RNA	2511 87384281 (5021, 5022)	2512   88084771 (5023, 5024)				2516 87786908 (5031, 5032)	2517 87784966 (5033, 5034)

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NCLASSIFIED	kinase transport	histone	transport
Contains protein domain (PF00018) - UNCLASSIFIED SH3 domain		Contains protein domain (PF00850) - histone Histone deacetylase family	Contains prolein domain (PF01753) - MYND finger
1jAF15181 - (AF151819) CGI-61	100 2	gil4826722[ref]NP_005085.1 pFATP - fatty acid transport protein 4 Novel Protein sim. GBank Novel Protein sim. GBank gil5174489[ref]NP_006035.1 pKlAA - histone deacetylase 6 Histone deacetylase family	87754052 (5043, 5044) Novel Protein sim. GBank glk580011[gblAAD24201.1[U81002 (U81002) TRAF4 associated factor 1 [Homo sapiens] 95340467 (5045, 5048) 95340469 (5047, 5048) BOP2 [Mus musculus]
2518   84147410 (5035, 5036)   Novel Protein sim. GBank   8149285919b AAD34056,   protein [Homo sapiens]	2519   94326180 (5037, 5038)   Novel Protein sim. GBank (AC004883) similar to KIA (AC004883) sim	gil4826722[ref]NP_005085 2521 95316244 (5041, 5042) Novel Protein sim. GBank gil5174489[ref]NP_006035	2522 87754052 (5043, 5044) ( 2523 95340467 (5045, 5048) 2524 95340469 (5047, 5048) (

				-	
\$ 525	94126928 (5049, 5050)	5525 94126928 (5049, 5050) Novel Protein sim. GBank gi[2073564 (U80223) - eukaryotic initiation factor elF-2 alpha kinase; DGCN2 [Drosophila melanogaster]		kinase	264488, 22278997, 22278999, 60432049, 60432289, 29331828, 264905, 265008, 55812038, 21906754, 265019, 264369, 21906765, 21906766, 21906767, 21908769, 35695917, 265020, 265021, 33657109, 60431528, 83373044, 60432113, 22279000,
<del></del>	95289404 (5051, 5052)	ile [Homo saplens]	Contains protein domain (PF00238) - ribosomalprot Ribosomal protein L14		60424179, 264768, 264687, 264769, 264689, 65274572, 21906767, 56182575, 21906768, 21906768, 22278994, 22278995, 3569528, 25278995, 3569528, 25278994, 22278995, 3569520, 22278994, 22278997, 265020, 22278998, 265021, 22278999, 265022, 264690, 264691, 60432049, 264259, 264097, 33657023, 29331827, 29331828, 246692, 29331827, 29331828, 246692, 29331827, 29331828, 246907, 18108370, 66712502, 60431528, 264907, 18108372, 18108374, 56182435, 18108376, 55811578, 35696423, 35695855, 265006, 265007, 264518, 264595, 264595, 60431735, 264638, 264595, 5581238, 60433356, 6043348, 264595, 5581238, 264596, 264598, 264598, 264598, 264598, 264598, 264513, 265018, 265018, 264586, 264587, 264567, 22279000, 265018, 264588, 264587, 264567, 264566, 264588, 264587, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264568, 264567, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264568, 264567, 264567, 264568, 264567, 264567, 264568, 264568, 264568, 264567, 264567, 264568, 264568, 264567,
2527	88094580 (5053, 5054)	gi]2258437 (AF008197) - us]		UNCLASSIFIED	264510, 264512, 264630, 264591, 264592, 264259, 284594, 264595, 264603, 264605, 18108351, 264565, 284369, 18108354
			Contains protein domain (PF00651) - dna_ma_bind BTB/POZ domain	dna_ma_bind	55812038, 265017, 264689, 35695917, 35695763, 60431528, 60432113, 22279002
2529			,	synthase	264908, 264769, 265020, 265021, 18108383
2530	80259978 (5059, 5060)				264369, 264556
2531				UNCLASSIFIED	29331822, 29331824, 60432289, 264508, 264509, 264906, 265011, 264769, 21906768. 33657023, 87168518, 22279000
2532		87419778 (5083, 5064) Novel Protein sim. GBank gi 2864625 emb CAA16972  -  (AL021811) putative protein (Arabidopsis thaliana)			284593
2533	87000255 (5065, 5066	87000255 (5065, 5066) Novel Protein sim. GBank gil437181 (U02289) - GTPase- activating protein [Caenorhabditis elegans]		UNCLASSIFIED	264555

D 264259, 35696052, 264905, 265017, 21906769, 265020, 265022, 33657109, 22279000	65274572, 35696286, 60432289, 29331828, 66712502, 265008, 60432229, 265017, 265018, 265019, 264288, 264369, 264689, 21906768, 265020, 265021, 264636, 60170394, 22279002	18108398, 56182575, 35696286, 22278997, 22278999, 60432049, 264259, 29331824, 29331826, 29331827, 29331827, 29331827, 29331827, 29331828, 264695, 264591, 264596, 52646317, 18108351, 264681, 264683, 18108354, 264289, 21906766, 21906767, 265021, 52645129, 33657409, 18108388, 87168518, 60432113, 22279000, 22279002, 264567, 18108391	65274572, 35696286, 29331822, 29331825, 29331827, 29331828, 35696052, 264906, 66712502, 264909, 265008, 265011, 264760, 264288, 264685, 35695917, 60170615, 264691, 33657023, 65274620, 33657109, 18108374, 35696423, 35695855, 264636, 264558, 60170394, 56182323, 83373044		ED 22278996, 22278999, 29331822, 29331825, 29331828, 29146499, 264908, 264112, 60170831, 87168559, 264604, 265019, 264685, 264766, 87168518, 22279000, 264565, 264566		ED 264488, 264768, 264689, 264511, 20281171, 264634, 264635, 264691, 264603, 264604, 264905, 264907, 264908, 704768
UNCLASSIFIED		kinase	ubiquitin	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED
			,				
2534 87332322 (5067, 5068) Novel Protein sim. GBank gij3452473 (AF084205) - serine/threonine protein kinase TAO1 [Rattus norvegicus]	91225056 (5069, 5070) Novel Protein sim. GBank gil4468311 emb CAB37992  - (AL031432) dJ465N24.2.1 (PUTATIVE novel protein) (isoform 1) [Homo saplens]	94218540 (5071, 5072) Novel Protein sim. GBank gij728836jspjP39193jALUB_HUMAN - IIII ALU SUBFAMILY SP WARNING ENTRY IIII	95422283 (5073, 5074) Novel Protein sim. GBank gil4557026jref]NP_003913.1 pHERC - guanine nucleotide exchange factor p532			2540   94218545 (5079, 5080) Novel Protein sim, GBank gil1362647 pir  S53876 - sex- regulated protein janus A - fruit fly (Drosophila pseudoobscura)	95308238 (5081, 5082) Novel Protein sim. GBank gi 1711658 sp P54797  T10_MOUSE - SER/THR-RICH PROTEIN T10 IN DGCR REGION
87332322 (5067, 5068)		94218540 (5071, 5072)		36853454 (5075, 5076)	94144916 (5077, 5078)	84218545 (5079, 5080)	
2534	2535	2536	2537	2538	2539	2540	2541

22	95298162 (5083, 5084)		Contains protein domain (PF00220) - UNCLASSIFIED		264488, 18108394, 52646385, 52646842,
		gripzzpazdigetAALAugountAFosaru - (AFosarun) sinuin	Neuronypopmysial normones, it-		05274374, £2276334, 03030200, £2270330, 054259 52645080 29331822 29331824.
_		type 2 [rioing septems]			29331827, 35696052, 33656970, 264907,
					264909, 52644045, 264510, 265008, 264512.
				•	265007, 265008, 265009, 264910, 60431735,
					52646317, 52644296, 265010, 265011,
					265018, 265019, 18108351, 264683, 264288,
					264685, 264687, 52644229, 264769,
					21906766, 21906767, 21906769, 52644150.
					33657023, 33657109, 52645129, 33657182,
					27486261, 27486264, 33657349, 35695763,
					18108374, 35696423, 35695855, 264631,
			,		264634, 264635, 264558, 83373044,
					18108385, 18108387, 87168518, 264563.
					264564
2543	94139088 (5085, 5086)	94139088 (5085, 5086) Novel Protein sim. GBank gij5419857 emb CAB46374.1 -	Contains protein domain (PF00076) -		65274572, 56182575, 22278999, 264259,
		(AL096723) hypothetical protein [Homo saplens]	RNA recognition motif. (a.k.a. RRM.		29331826, 264907, 264510, 264511, 264592.
			RBD, or RNP domain)		264595, 264764, 264369, 264288, 264684,
					264766, 264689, 21906765, 21906767,
					21906769, 60170615, 264692, 264693,
_					55811576, 65274791, 264636, 264556,
					18108381, 60170394, 264639, 18108385,
					60432113, 22279000
2544	94218549 (5087, 5088)	94218549 (5087, 5088) Novel Protein sim. GBank	Contains protein domain (PF00629) - glycoprotein		18108397, 52646365, 22278997, 264259.
		gij2498110jspjQ63191jAEGP_RAT - APICAL ENDOSOMAL	L MAM domain.		60432049, 29331822, 29331825, 29331826.
		GLYCOPROTEIN PRECURSOR			29331827, 29331828, 264905, 264908,
					265006, 265007, 265008, 87168559, 265017,
					265018, 265019, 18108351, 264448, 264686,
	•				264687, 264689, 21906765, 265020, 265021.
					18108370, 18108374, 18108376, 18108381,
					18108385, 18108387, 56526486, 22279000,
					264482, 264563, 264567
¥5	87742645 (5089, 5090)	2545  87742645 (5089, 5090)   Novel Protein sim. GBank gij3327046 dbj BAA31591  -			29331825, 264906, 265009, 60170831,
		(AB014516) KIAA0616 protein [Homo sapiens]			265017, 264369, 21906767, 60170615,
1				G11004 10111	204034, 33037 103
2746		88093861 (5091, 5092) Novel Protein sim. GBank gij2996032 (AF054586) - brain	Contains protein domain (PF00097) - UNCLASSIFIED	UNCLASSIFIED	29331624, 265001, 22219002
		Inger protein [Raffus norvegicus]	Zinc finger, C3HC4 type (RING		
			linger)		

264488, 18108394, 52646842, 18108397, 56182575, 22278995, 56994075, 22278996, 22278997, 22278995, 56994075, 22278996, 22278997, 22278999, 264259, 29331822, 29331824, 29331826, 60432289, 29331827, 35696052, 264308, 66712502, 264509, 264905, 264906, 264906, 264908, 66712502, 264909, 265009, 60170831, 60432229, 60431735, 265009, 60170831, 60432229, 60431735, 265019, 265011, 264600, 264601, 265018, 265019, 264760, 18108351, 264682, 264488, 264288, 264369, 264684, 264682, 264697, 265019, 265019, 264691, 18108362, 265021, 265022, 264690, 264631, 18108362, 264692, 264693, 264691, 18108362, 264692, 264693, 264631, 18108362, 264692, 264693, 264631, 18108370, 18108370, 264692, 264693, 264638, 56182323, 264558, 264556, 264566, 264566, 264561, 18108386, 264556, 264566, 264566, 264561, 18108385, 264558, 264566, 264566, 264566, 264569, 264561, 18108385, 264558, 264566, 26	264489. 18108394, 52646365, 22278994, 25686286, 56994075, 22278997, 22278999, 264259, 29331822, 29331827, 29331825, 29331826, 29331826, 29331827, 29331827, 29331828, 56182435, 264511, 265007, 264512, 60433356, 87168559, 264684, 264369, 52644229, 265021, 33657023, 264692, 18108374, 52644332, 264557, 18108381, 18108382, 18108381, 18108382, 18108382, 264563, 264567	56182575, 22278936, 22278937, 22278938, 22278939, 22278939, 264259, 264508, 264908, 29331839, 265009, 265010, 265018, 264688, 21906769, 21906766, 21906766, 21906769, 265020, 265021, 52644150, 264691, 18108368, 60431602, 18108376, 35696423, 56182323, 18108387, 264567	56182575, 29331822, 264105, 264512, 18108351, 35695917, 264637, 264638
ransport	UNCLASSIFIED	ਜ਼ਹੂਰ -	UNCLASSIFIED
Mitochondrial carrier proteins Mitochondrial carrier proteins		Contains protein domain (PF00412) - struct LIM domain containing proteins	
2547 94143869 (5093, 5094) Novel Protein sim. GBank gil4929607[gblAAD34064.1 AF15182 - (AF151827) CGI-69   protein [Homo sapiens]	6	94196893 (5097, 5098) Novel Protein sim. GBank gij728837 sp P39194 b4LU7_HUMAN - IIII ALU SUBFAMILY   LIM domain containing proteins SQ WARNING ENTRY IIII	87778584 (5099, 5100) Novel Protein sim. GBank gij2143886 pir  i52523 - Inucleoporin p62 homolog - ral (fragment)
3. 5094)	5096)	97, 5098)	99, 5100
94143869 (506		2549   94196893 (50)	87778584 (50

95309400 (5101, 5102) Novel Protein sim, CBank gil4337103igbjAAD18079] - Amadilabeta hydrolase fold (AF129789) NG26 [Homo sapiens]   Amadilabeta hydrolase fold   95309202 (5103, 5104)   Novel Protein sim, CBank gil4331 (99092031   EBM-1   LUNCLASSIFIED   95309203 (5103, 5105) Novel Protein sim, CBank gil4331 (99092031   EBM-1   LUNCLASSIFIED   95309203 (5110, 5105) Novel Protein sim, CBank gil4331 (99092031   EBM-1   LUNCLASSIFIED   95309203 (5111, 5110) Novel Protein sim, CBank gil43399991emb[CA803030] - Amadilabeta-catenin-like repeats (ALGO0904) photolerial min, CBank gil4339991emb[CA803030] - Amadilabeta-catenin-like repeats (ALGO0904) photolerial min, CBank gil4339991emb[CA803030] - Amadilabeta-catenin-like repeats (ALGO0904) photolerial min, CBank gil4339991emb[CA803030] - Amadilabeta-catenin-like repeats (ALGO0904) photolerial min, CBank gil4339991emb[CA803031] - Contains protein domain (PF00514) - UNCLASSIFIED (ALGO803) 57341.3 (min) (olidadoy receptor ULE) (hode pain family) (olidadoy receptor ULE) (hode pain family) (hode Protein sim, CBank gil533991emb[CA803031] (hode Protein sim, CBank gil533991emb[CA803031] (hode Protein sim, CBank gil533991emb[CA803031] (hode Protein sim, CBank gil43399) (hode Protein sim, CBank gil533991emb[CA803031] (hode Protein sim, CBank gil6313991emb[CA803031] (hode Protein sim, CBank gil63139391emb[CA803031] (hode Protein sim, CBank gil63139391emb[CA80303031] (hode Protein sim, CBank gil63139391emb[CA8030303031] (hode Protein sim, CBank gil63139391emb[CA803030303031] (hode Protein sim, CBa	22278998, 22278999, 264259, 29331822, 29331824, 29331824, 29331826, 29331827, 29331828, 265007, 60432229, 87168559, 265017, 265018, 265019, 264689, 21906766, 21906769, 35695917, 265020, 33657023, 33657109, 18108374, 264634, 264559, 18108385, 87168518, 22279002		22278997, 29331822, 29331828, 60433356, 265011, 264288, 264765, 264766, 264769, 21906765, 21906766, 60432113, 264482	264906, 264511, 265018, 264448	264595	264684, 264766, 264689, 33657023, 33657109, 35695855, 264558, 264567	29331824, 263972	22278997, 29331822, 264508, 21908769, 33657023, 33657109, 56182323	264762, 264906, 264592, 264901, 264566, 264908, 264684, 264567, 264909, 264766	264686, 26448, 263976, 264768, 29331626, 35696052, 35696423, 264601, 264511, 264602, 264910, 264634, 264760, 264555,	33657109, 18108374, 55811576, 35696423, 35695855, 56182323, 264558	56182435, 265008, 264591, 33109954, 264760, 55811957, 35695917, 33657023,	56182575, 35696286, 29331824, 29331826, 35696052, 29331828, 264508, 264907,	56182323, 83373044, 22279000	21906768, 21906769, 265020, 60170615,	265018, 264605, 265019, 264288, 21906766,	55812038, 21906754, 65274444, 265017,	29331825, 60432289, 29331826, 29331827.	22278999, 264259, 29331822, 29331824,	118108396 65274572 22278995 22278998
9[4337103]gb[AAD18079] - 6 saplens]  AEGION  EGION  - MOUSE - SER/THR-RICH  EGION  - HUMAN - IIII ALU SUBFAMILY  gi]4884319]emb[CAB43260.1] - 1000  gi]4106984 (AC003038) - 11  gi]4538999]emb[CAB44995.1] - 117M1 - 4 (novel 7 transmembrane 1) (olfactory receptor LIKE)  C.1 JAF04495 - (AF044953)  sductase PGIV subunit [Homo	·		UNCLASSIFIED				nuclease	cadherin		UNCLASSIFIED			UNCLASSIFIED			Š				INCI ASSIFIED
gjl4337103jgbjAAD18079j - saplens]  MOUSE - SER/THR-RICH REGION  E-HUMAN - III! ALU SUBFAMILY gjl4884319jembjCAB43260.1j - rotein [Homo saplens] gjl4106984 (AC003038) - sij119110jspjP03211jEBN1_EBV TEIN gjl453899gjembjCAB4995.1j - tein [Arabidopsis thaliana] tein [Arabidopsis thaliana] sgjl5051399jembjCAB4995.1j - tein [Arabidopsis thaliana] sgjl45081999gjembjCAB4995.1j - tein [Arabidopsis thaliana]	·	Contains protein domain (PF00001) - 7 transmembrane receptor (rhodopsin family)				Armadillo/beta-catenin-like repeats													alpha/beta hydrolase fold	Contains profein domain (PE00561) -
0 (5103, 5102 0 (5103, 5106 3 (5105, 5106 11 (5109, 5110 13 (5111, 5112 14 (5113, 5114 16 (5119, 5120 15 (5117, 5112	gij5326825[gbJAAD42056.1AF04495 - (AF044953) NADH:ubiquinone oxidoreductase PGIV subunit [Homo sapiens]	) Novel Protein sim. GBank gil5051399Jemb[CAB44995.1] - (AL078630) 573K1.3 (mm17M1-4 (novel 7 transmembrane receptor (rhodopsin family) (olfactory receptor LIKE) protein)) [Mus musculus]	) Novel Protein sim. GBank gil4538998 emb CAB39619.1  - (AL049481) AIG1-like protein [Arabidopsis thaliana]	- EBNA-1 NUCLEAR PROTEIN	(1	R30923_1 [Homo sapiens]	Novel Protein sim. GBank gij4884319jemb CAB43260.1  - (AL050084) hypothetical protein [Homo sapiens]	) Novel Protein sim. GBank gi 728835 sp P39192 ALU5_HUMAN - IIII ALU SUBFAMILY SC WARNING ENTRY IIII		~ ~			10							
2551 95308400 2552 9533262 2553 9530824 2554 8776152 2555 8762755 2556 8764553 2556 87843786 2557 7843786 2557 7843786 2556 8789453				_									95332620 (5103, 5104)						(1000) 000000	195308400 (5104 5102)

56994075, 22278996, 22278997, 22278999, 264259, 29331822, 60432289, 33657402, 60433356, 21908765, 55811957, 60170615, 33657023, 264693, 35695855, 87168518	264488, 35696286, 22278999, 264259, 29331822, 29331824, 35696052, 264508, 264907, 264908, 264909, 52644045, 264510, 264511, 265009, 264910, 264591, 265411957, 3656019, 18108351, 264686, 21906767, 21906768, 55811957, 35695917, 265020, 264691, 264693, 256693, 264632, 18108374, 264632, 264632, 18108384, 87168518, 22279000, 22279002, 264486	29331822, 265007, 265010, 265019, 264769, 55811576, 56182323	66714117, 264909, 263978, 264632	18108370, 35695855, 264556, 264558, 18108383	265020, 60170615	60424179, 18108394, 56181686, 56994075, 2227899, 264490, 264259, 29331822, 56182181, 29331824, 60424269, 29331825, 29331826, 29331826, 29331826, 29331826, 29331826, 265009, 33657402, 265010, 265011, 265019, 264448, 264683, 264288, 18108354, 264769, 21906766, 25659517, 265021, 33657023, 1810836, 33657109, 33657182, 35695763, 60431528, 55810764, 18108379, 83373044, 18108385, 60432113, 264482
UNCLASSIFIED	UNCLASSIFIED	sulfotransferase	UNCLASSIFIED	struct		synthase
:				Contains protein domain (PF00063) - struct Myosin head (motor domain)		
2562   87645539 (5123, 5124) Novel Protein sim. GBank gil4106984 (AC003038) - R30923_1 [Homo sapiens]	88095497 (5125, 5126) Novel Protein sim. GBank gil4886447 emb CAB43371.1  - (AL050270) hypothetical protein [Homo sapiens]	80502783 (5127, 5128) Novel Protein sim. GBank gil 352944 sp P47179 YJ9P_YEAST - HYPOTHETICAL 118.4 KD PROTEIN IN BAT2-DAL5 INTERGENIC REGION PRECURSOR	130)	80224956 (5131, 5132) Novel Protein sim. GBank gil628012 pir  A53933 - myosin l   myr 4 - ral	134)	91233099 (5135, 5136) Novel Protein sim. GBank gil466009lsplP34548 YNJ4_CAEEL - HYPOTHETICAL 33.8 KD PROTEIN R10E11.4 IN CHROMOSOME III
87645539 (5123, 512	88095497 (5125, 512		ı			
2562	2563	2564	2565	2566	2567	2568

18108394, 56182575, 56181686, 22278995, 22278996, 56994075, 35696286, 22278997, 22278998, 264259, 60432049, 22378999, 264259, 60432049, 29331824, 29331824, 29331825, 60432289, 29331826, 29331827, 29331825, 265008, 264906, 5618243, 265008, 265009, 264906, 5618243, 265017, 265018, 265019, 264682, 264448, 264288, 21906765, 21906768, 21906767, 21906768, 29148627, 21906769, 35695917, 265021, 265022, 2564150, 264690, 264691, 265022, 264691, 265022, 264691, 265022, 25644150, 264690, 264691, 265022, 2646150, 263655, 264691, 265022, 26461069, 263974, 18108374, 55810764, 35696423, 35695655, 264558, 18108385, 56526468, 87168518, 22279000, 22279002, 284566	22278996, 29331822, 29331824, 66714117, 29331825, 60432289, 29331827, 35696052, 264907, 264510, 265007, 265009, 264758, 33109954, 265019, 264686, 264689, 265020, 265021, 264691, 264692, 264693, 18108374, 35695855, 284634, 56182323, 264639, 60170394, 83373044, 22278002, 264482	264558, 264905, 264907, 264828, 18108351, 264555, 264556, 264557, 264558, 264559	35696286, 29331827, 35696052, 264100, 264104, 264110, 264592, 21906754, 29148627, 29148629, 263972, 263974, 18108374, 263976, 35695855, 60170394, 264559, 18108385,
transcriptfactor	phosphatase	struct	UNCLASSIFIED
Contains protein domain (PF00170) - transcriptfactor	Contains protein domain (PF01240) - phosphatase Protein phosphatase 2A regulatory subunit PR55		
2569   95313764 (5137, 5138) Novei Protein sim. GBank gi 2599560 gb AAB84166.1  - (AF029674) basic leucine zipper protein LZIP [Homo sapiens]	2570 94136754 (5139, 5140) Novel Protein sim. GBank gil4758954 ref NP_004567.1 pPPP2 - protein phosphalase 2 (formerly 2A), regulatory subunit B (PR 52), beta isoform	87733750 (5141, 5142) Novel Protein sim. GBank gij732218 sp P34609 YO60_CAEEL - HYPOTHETICAL 128.6 KD PROTEIN ZK1098.10 IN CHROMOSOME III	2572   87627560 (5143, 5144) Novel Protein sim. GBank gil4884319jemb CAB43260.1
N (5137, 5138) N (5137, 5138) N (5137, 5138)	94136754 (5139, 5140) N	87733750 (5141, 5142) h	8,7627560 (5143, 5144) h
8999	2570	2571	2572

2573	95313929 (5145, 5146)		Contains protein domain (PF00386) - complement		264488, 60424179, 65274572, 56182575,
		gij399138jsp[P02745]C1QA_HUMAN - COMPLEMENT C1Q C1q domain SUBCOMPONENT, A CHAIN PRECURSOR	C1q domain	2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	260181086, 22278935, 20334824, 20331824, 20331828, 20331822, 20331827, 20331827, 20331828, 20331827, 20331828, 20331828, 20331827, 20331828, 20331827, 20331828, 20331827, 20331827, 20331827, 20331827, 20331830, 264906, 204310, 264901, 264906, 264909, 264910, 264591, 264592, 60432229, 264593, 60433356, 264594, 60433438, 264595, 55612038, 264759, 264077, 265019, 265011, 264761, 264761, 264762, 264763, 264764, 264869, 264763, 264769, 264763, 26489, 21906766, 21906766, 2918677, 264689, 21906768, 21906769, 265021, 265021, 265021, 264691, 264692, 33657023, 657747610, 264691, 264692, 33657023, 657747610, 264691, 264632, 264639, 21908768, 21908344, 21908348, 21908348, 21908768, 21908344, 21908348, 21908768, 21908768, 21908768, 21908344, 21908768, 21908344, 21908348, 21908768, 21908344, 21908348, 21908768, 21908344, 21908348, 21908768, 21908344, 21908348, 21908768, 21908768, 21908344, 21908348, 21908768, 21908344, 21908344, 21908768, 2190876
2574	94746814 (5147, 5148)	94746814 (5147, 5148) Novel Protein sim. GBank gij3334982 (AC005306) - R27216_1 [Homo sapiens]	Contains protein domain (PF00651) - UNCLASSIFIED BTB/POZ domain		22278995, 264259, 60432289, 29331827, 29331828, 33656970, 264909, 265008, 265619, 264410, 264591, 33657402, 265018, 26519, 264448, 264764, 264369, 24508768, 18108357, 21906765, 21906768, 55811957, 60170615, 264691, 33657023, 264693, 33557109, 33657182, 27486264, 33567349, 264636, 264555, 83373044, 18108385, 264482
2575		87754408 (5149, 5150) Novel Protein sim. GBank gif4929729[gbJAAD34125.1]AF15188 - (AF151888) CGI-130 protein [Homo sapiens]		UNCLASSIFIED	264910, 264601, 264683, 264689, 264080
2576		95357881 (5151, 5152) Novel Protein sim. GBank gil4680681lgblAAD27730.1JAF13295 - (AF132955) CGI-21 protein [Homo sapiens]	Contains protein domain (PF00442) - ubiquitin Ubiquitin carboxyl-terminal hydrolases family 2		264259, 35696052, 264906, 60433438, 264681, 18108351, 264288, 52644150, 264628, 35696423
2577	86996621 (5153, 5154)	86996621 (5153, 5154) Novel Protein sim. GBank gild337103lgb AAD18079  - (AF129756) NG26 [Homo saplens]		UNCLASSIFIED	29331825, 265018, 265019, 264685
2578	87786941 (5155, 5156)			UNCLASSIFIED	264488, 264906, 264908, 264910, 264596, 264603, 264604, 264605, 264768, 21906769, 264628, 264630, 264634, 264639, 264563
2579	87292879 (5157, 5158)			UNCLASSIFIED	29331822, 29331824, 264767

265007, 265018, 264762	56994075, 29331824, 29331826, 29331828, 264905, 60433356, 60433438, 264758, 87168559, 21906769, 265022, 35695855, 263981	264488, 264907, 264908, 264910, 264764, 264684, 264766, 264636, 264555, 264565	56181686, 264259, 264510, 264512, 264591, 264592, 264592, 264593, 264594, 284595, 264596, 264603, 264629, 55810764, 264630, 264637, 264565	264908, 264910, 264768, 264693, 18108374, 55811576, 56182323	264768	22278998, 264259, 29331822, 29331824, 29331827, 29331827, 29331827, 265007, 265009, 264591, 60433356, 33657402, 265018, 264762, 264288, 21906766, 21906767, 21906769, 265202, 264691, 83373044, 56528486, 22279002	284908, 265019, 264768, 264693, 55811576, 56182323	264564	256909, 265009, 265018, 264769, 35696423, 264636	[22278995, 29331830, 265008, 265010, 265017, 264639	264259, 29331822, 264106, 264906, 56182435, 265007, 265008, 33109954, 264448, 55811957, 265020, 18108370, 55811576, 22279002
	collagen	eph	transcriptfactor	UNCLASSIFIED	kinase			UNCLASSIFIED	ONCEASSIFIED	glucoamylase	eph
	Contains protein domain (PF00595) - collagen PDZ domain (Also known as DHR or GLGF).		Contains protein domain (PF00047) - Irranscriptfactor Immunoglobulin domain							Contains protein domain (PF01055) - glucoamylase Glycosyl hydrolases family 31	
Novel Protein slm. GBank gi[2588628 (AC003080) - Similar Io KIAA0299: 60% slmilarity to AB002297 (PID:92224539) Homo sapiens]	Novel Protein sim. GBank gil4406642 gb AAD20049  - (AF131809) Unknown [Homo sapiens]	Novel Protein sim. GBank gil2739367 (AC002505) - putative phosphatidylinositol-4-phosphate 5-kinase (Arabidopsis thaliana)	Novel Protein sim. GBank gi 4378112 emb CAA16521.1  - Contains protein domain (AL021578) dJ453C12.2 (similar to transcription factor RBP- Immunoglobulin domain L) [Homo sapiens]		80436126 (5169, 5170) Novel Protein sim. GBank gi[2736151 (AF021935) - mytonic dystrophy kinase-related Cdc42-binding kinase [Rattus norvegicus]			13777E37601 1003760001 10037	65515607 (5177, 5178) Nover Protein sim. Gbank gij3021398jembjCAA71415j - (Y10389) nuclear protein [Xenopus laevis]	87054528 (5179, 5180) Novel Protein sim. GBank gij2104689 (U92793) - alpha glucosidase II, alpha subunit [Mus musculus]	94192167 (5181, 5182) Novel Protein sim. GBank gi 5702202 gb AAD47199.1 AF12916 - (AF129166) long- chain acyl-CoA synthetase 5 [Homo sapiens]
2580 88166788 (5159, 5160) Novel Protein sim. GBank to KIAA0299; 60% similari [Homo sapiens]	87899048 (5161, 5162) Novel Protein sim. GBank (AF131809) Unknown [Ho	2582 87786789 (5163, 5164) Novel Protein sim. GBank phosphatidylinositol-4-pho (thallana)	2583   91220950 (5165, 5166)   Novel Protein sim. GBank (AL021578) dJ453C12.2 ( L) [Homo sapiens]	80430941 (5167, 5168)	80436126 (5169, 5170) h	2586 91226136 (5171, 5172)	80430943 (5173, 5174)	80074385 (5175, 5176)	(8716,7716) //Joc 1668	87054526 (5179, 5180) I	94192167 (5181, 5182)
2580	2581	2582	2583	2584	2585	2586	2587	2588	8907	2590	2591

WO 00/58473 PCT/US00/08621

292	95332648 (5183, 5184)	2592   95332648 (5183, 5184) Novel Protein sim. GBank gi]3024998 sp Q60936 YAB1_MOUSE - HYPOTHETICAL HEART PROTEIN		sport	18108397, 56182575, 35696286, 56994075, 284259, 29331824, 29331828, 66932299, 29331824, 29331828, 29331828, 284206, 284599, 28331827, 284390, 60432229, 284594, 60433356, 60433438, 55812038, 18108348, 21906754, 265011, 87168559, 265017, 265019, 284764, 26399, 264288, 264766, 265021, 60170615, 33657023, 33657109, 264629, 35696423, 35695855, 264557, 264638, 60170394, 56182323, 264482, 264565
	87754416 (5185, 5186)	87754416 (5185, 5186) Novel Protein sim. GBank gil4929729[gb AAD34125.1 AF15188 - (AF151888) CGI-130 protein [Homo sapiens]			22278999, 29331825, 264758, 21906754, 52646317, 265010, 18108351, 264288, 264369, 21906768, 264693, 18108370, 264638, 264482
2594		95305758 (5187, 5188) Novel Protein sim. GBank gil4929587Igb AAD34054.1 AF15181 · (AF151817) CGI-59 protein [Homo sapiens]		UNCLASSIFIED	264488, 18108398, 56182575, 35696286, 2278997, 264093, 264259, 29331822. 29331825, 264093, 264259, 29331822. 29331825, 264905, 264909, 5264009, 5264204, 264510, 264512, 265007, 264757, 21908754, 264764, 264760, 264448, 264764, 264288, 21906768, 33657109, 263997, 264634, 264586, 264567, 264564, 264567, 264567
2595	į			UNCLASSIFIED	264692
2596		87538637 (5191, 5192) Novel Protein sim. GBank gil4309681 gb AAD15478 - , (AC006930) R33423_1 [Homo sapiens]		UNCLASSIFIED	22278999, 264259, 265018, 264448, 265021, 60431528
2597	94784089 (5193, 5194)			UNCLASSIFIED	264905, 264509, 264908, 264766, 35695917, 35695855, 264635, 264636. 83373044, 264486
2598		88094948 (5195, 5196) Nover Protein sim. GBank gij1001351jdbjjBAA10838j - (D64006) hypothelical protein [Synechocystis sp.]		UNCLASSIFIED	22278998, 264259, 29331824, 87168474, 264683, 21906766, 35695917, 264691, 33657023, 33657109, 18108370, 18108374, 264564, 264565
2599		87642889 (5197, 5198) Novel Protein sim. GBank gij3941737 (AF109719) - BAT2 [Mus musculus]		МНС	264766, 264769, 21906768, 33657182, 35695763, 18108370, 18108374, 264635, 264636, 56528486, 22279000, 264566
2600		87787846 (5199, 5200) Novet Protein sim. GBank gil4263521gb AAD15347  - (AC004044) putative WD-repeat protein (Arabidopsis thaliana)	Contains protein domain (PF00400) - kinasereceptor WD domain, G-beta repeat	kinasereceptor	35696286, 264093, 264288, 21906769, 35696423, 35695855

29331822, 29331825, 29331826, 29331827, 29331828, 264908, 264907, 264908, 66712502, 264828, 56182435, 55812038, 265010, 265017, 265018, 26509, 264689, 21906765, 55811957, 285020, 265022, 264692, 33657023, 264693, 33657109, 18108370, 264639, 565182	264509, 264907, 264908, 264592, 264755, 264631	224488, 65274572, 35696286, 22278997, 22278999, 60432049, 264259, 56182181, 29331824, 29331826, 60432289, 29331824, 29331826, 60432289, 6043336, 60433438, 55812038, 264907, 60433438, 55812038, 26448, 264765, 264288, 264766, 264689, 21906765, 21906767, 21906769, 265017, 26493, 24692, 33657109, 27486261, 18108370, 65274791, 264639, 264556, 264533, 38168385, 56526486	264259, 2231822, 2931825, 60432289, 29343827, 264107, 264109, 264905, 26182435, 264112, 265008, 265007, 265008, 265009, 60433356, 60433438, 265011, 87168559, 265017, 264448, 264682, 264764, 264288, 265021, 33657023, 263967, 33657182, 27486261, 18108374, 263976, 55811576, 264638, 87168518, 60432113	22278999, 265017, 264684, 21906768, 22278000 22278000 265009, 264910, 264759, 265017, 21906767, 18108365, 18108388, 60432113	264685 22278998, 22278999, 29331825, 264508, 264906, 21906754, 264602, 264766, 264769, 52644229, 21906765, 33657109, 27486264, 18108370, 263972, 284555, 60432113	264768, 18108394, 264692, 264693, 264509, 264509, 264807, 284628, 284908, 264909, 18108377, 284511, 284512, 264910, 264635, 264595, 265010, 264404, 264563, 264764, 264685, 264766
		histone	ranscription of the control of the c	isomerase Iransport	-	UNCLASSIFIED
		Contains protein domain (PF00850) - histone Histone deacetylase family		Contains protein domain (PF00254) - isomerase FKBP-type peptidyl-prolyl cis-trans isomerases transport	Contains protein domain (PF00625) Guanylate kinase	
87627742 (5213,.5214) Novel Protein sim. GBank gil4826626[gb[AAD30202.1] - (AF135022) mediator [Homo saplens]	81734786 (5215, 5216) Novel Protein sim. GBank gij2226005 (U49973) - ORF2: function unknown [Homo sapiens]	94843791 (5217, 5218) Novel Protein sim. GBank gil3024889 sp P56524 Y288_HUMAN - HYPOTHETICAL PROTEIN KIAA0288 (HA6116)	88177654 (5219, 5220) Novel Protein sim. GBank gil4336855 gb AAD17889] - (AF106473) leucine-rich-domain inter-acting protein 1; LEAP1 [Mus musculus] inter-acting protein 1; LEAP1 [Mus musculus]	87428890 (5221, 5222) Novel Protein sim. GBank gij3876761[emb]CA492994] - Contains protein (268760) predicted using Genefinder; Similarity to Mouse FKBP-type protein (SW:FKB3_MOUSE) (Caenorhabditts isomerases elegans)   Regional   Regional	79481496 (5225, 5226) 87643948 (5227, 5228) Novel Protein sim. GBank gij5533081jgbjAAD45009.1jAF16118 - (AF161181) P55T protein [Mus musculus]	
87627742 (5213,.5214)	81734786 (5215, 5216) N	94843791 (5217, 5218) N	88177654 (5219, 5220)			87381996 (5229, 5230)
2607	2608	2609	2610	2611	2613	2615

2049. 11, 32,	11576,	64907. 768. 37.	54.	4572. 22. 1828. 1828. 6554. 64448. 44229. 109. 332.		512. 64636.	
22278995, 22278997, 22278998, 60432049, 60432288, 244828, 60433356, 264594, 60433438, 33109954, 87168474, 265011, 265019, 284288, 264768, 21906765, 21906767, 265020, 265021, 18108376, 18108377, 18108387, 87168518, 284482, 264567	265010, 265019, 264369, 264693, 55811576, 22279002	284489, 22278996, 264490, 264259, 29331822, 284102, 264509, 264908, 264907, 66712502, 29331830, 265008, 264908, 264907, 265009, 60433356, 60433438, 264758, 21906754, 265011, 87168559, 264768, 264689, 21906765, 21906765, 21906765, 21906767, 35659917, 265020, 265022, 33657023, 3657023, 3657023, 36595857, 18108381, 83373044, 18108385, 18108388, 56458481, 56458, 18108385, 18108388, 56458481, 56458, 18108385, 18108388, 56458481, 56458, 18108385, 18108388, 56458481, 56458, 18108385, 18108388, 56458481, 56458, 18108385, 18108388, 56458481, 56458, 18108385, 18108388, 56458481, 56458, 18108385, 18108385, 18108385, 18108385, 18108385, 18108388, 56458481, 56458, 18108385, 18108585, 181085	264259, 60432289, 60433438, 21906754, 264369, 60432113, 264566	5264507, 52645156, 52646842, 65274572. 22278995, 56994075, 35696286, 22278999, 60432049, 284259, 52645080, 29331822, 29331824, 29331825, 29331826, 29331826, 29331826, 29331827, 2859052, 264907, 66712502, 265008, 6043336, 33657402, 26546317, 21906754, 87168474, 265019, 265017, 265018, 264488, 264689, 21906765, 21906768, 35695917, 52644150, 264692, 33657109, 35695763, 35695713, 264362, 35677109, 35695763, 3695763, 3697727, 8716828, 86432713, 22779002, 5574727, 8716828, 86432713, 22779002, 5574727, 8716828, 86432713, 22779002, 56574727, 87168285, 86432713, 22779002, 56574727, 87168285, 87168385, 87168387, 8716837, 871687, 8716837, 8716837, 871687, 871687, 8716837, 871687, 871687, 871687, 871687, 871687, 871687, 871687, 871687, 871687, 871687,		264488, 264908, 264907, 264908, 264512, 265007, 264758, 35695917, 264634, 264636 264563, 264482	29331824, 35696052, 265007, 265010, 264288, 29148629
97, 22278, 6043335, 6043335, 87168, 28 84288, 26 75621, 87, 87168	64369, 26	284489, 22278996, 264490, 264259, 29331822, 264102, 264509, 264905, 265008, 264905, 265008, 264906, 265009, 265009, 265009, 60433356, 60433438, 26475, 21906754, 265011, 87168559, 26507, 266689, 21906765, 21906766, 21906766, 21906765, 21906765, 21906765, 21906766, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21806769, 264628, 181083, 26595855, 18108381, 83373044, 18	. 6043343 , 264566	56, 52646 75, 35696 25, 29331 . 6671250 02, 52646 . 265017, 1906765, 50, 26469 23, 26453		64907, 26 5695917,	52, 26500
5, 222789 9, 264828 3, 331099 265019, 2 7, 265020 7, 181083	265019, 2 2	22278996 2, 284102 2, 293318 60433356 60433356 4, 265011 264369, 2 21906765 7, 265020 33657109 8, 565784	264259, 60432289, 604334; 264369, 60432113, 264566	7, 526451 5, 569940 9, 264259 4, 293318 2, 264907 6, 336574 4, 265010 264289, 2 264689, 2 7, 526441 3, 356964 7, 871685	264636	264906, 2 264758, 3 264482	29331824, 356960: 264288, 29148629
2227899 6043228 6043343 265017, 2190676 1810837 264567	265010, 2 22279002	2933182. 2933182. 6871250. 265009. 2190675. 265618. 3569581. 264692.	264259. 264369.	5264450 2227899 6043204 2933182 3569605 8716847 8716847 264369 369591 3569591 3569591 1810838	264594, 264636	264488, 264906, 265007, 264758, 264563, 264482	2933182 264288,
		FIED .	nase	IFIED .			IFIED
somerase	kinase	UNCLASS	dehydroge	UNCLASS			UNCLASSIFIED
Contains protein domain (PF00254) - isomerase FKBP-type peptidyf-protyl cis-trans Isomerases		Contains protein domain (PF00397) - UNCLASSIFIED WW domain	Contains protein domain (PF00465) - dehydrogenase Iron-containing alcohol dehydrogenases	Contains protein domain (PF00628) - UNCLASSIFIED PHD-finger			
Contains protein domain (PF00254) FKBP-type peptidyf-prolyl cis-trans Isomerases		Iomain (PI	Jomain (Pi cohol	domain (Pl			5
s protein d rpe peptidi ses		nain nain	Contains protein domai Iron-containing alcohol dehydrogenases	s protein o			
Contains FKBP-ty s isomera		Contains pro	Contain Iron-con dehydro	Contains pr			
2994j • o Mouse norhabditk	HUMAN - IIII ALU SUBFAMILY	- (	putative nogaster]	-1260		Nover Frotein Sim. Obank gil4557341[ref]NP_001174.1[pATP6 - ATPase, H+ transporting. Iysosomal subunit 1; vacuolar proton pump; H- ATPase subunit	5299j - oditis
3876761 emb CAA92994 - enefinder, Similarity to Mous FKB3_MOUSE) {Caenomab	IIII ALU SI	AF049103 piens]	U66411) - hila melar	Novel Protein sim. GBank gild3225671gb AAD160 (AF090436) dachshund variant 1 [Mus musculus]		IlpATP6 - ATPase, H+ ıunit 1; vacuolar proton	ij3880355 emb CAB05299  enefinder [Caenorhabditis
1876761je nefinder; S KB3_MOL	- HUMAN	ij3319282 (AF046 in [Homo sapiens]	2431772 ( e [Drosop	nt 1 [Mus nt 1 [Mus		pATP6 - A nit 1; vacu	3880355 e nefinder [C
SBank gils using Ger in (SW:Fl	<b>-</b> '	SBank gil; ig protein	3Bank gil; drogenas	SBank gij	400	35871 001174.1  mal subur	SBank gil: using Ger
tein sim. C predicted iding profe	tein sim. (  sp P3918 NG ENTR	interactir	tein sim. cohot dehy	6) dachst	doin cim	I frefiNP og, lysoso ubunit	itein sim. predicted
Novel Protein sim. GBank gij3876/61 jemb CAA92994 - Contains pro (Z68760) predicted using Genefinder, Similarity to Mouse FKBP-type g FK508-binding protein (SW:FKB3_MOUSE) (Caenorhabdilis isomerases elegans)	Novel Protein sim. GBank gij728831 jspjP39188jALU1 J WARNING ENTRY III	Novel Protein sim. GBank g Huntingtin interacting protei	Novel Protein sim. GBank gi[2431772 (U66411) - putative type III alcohol dehydrogenase [Drosophila melanogaster]	Novel Pro (AF09043	Noted Dec	NOVEI FIGURI SIM. GDBMK gil4557341 prijNP_001174, transporting, lysosomal sub ATPase subunit	Novel Protein sim. GBank gij3880355 emb CAB05299 (282285) predicted using Genefinder [Caenorhabditis elegans]
1, 5232)	13, 5234)	5, 5236)	17, 5238)	19, 5240)	11. 5242)	3. 3244)	15, 5246)
AD 10 (87420895) (5231, 5232) Novel Protein sim. GBank gi (288760) predicted using Ge FK508-binding protein (SW: elegans)	86976888 (5233, 5234) Novel Protein sim. GBank  gil728831 sp P39188 ALU  J WARNING ENTRY !!!	91231662 (5235, 5236) Novel Protein sim. GBank gij3319282 (AF049103) Huntingtin interacting protein [Homo sapiens]	87694000 (5237, 5238) Novel Protein sim. GBank gi[2431772 (U66411) - putative type III alcohol dehydrogenase [Drosophila melanogaster]	95314841 (5239, 5240) Novel Protein sim. GBank gil4322567lgblAAD16097[ (AF090436) dachshund variant 1 [Mus musculus]	80253495 (5241, 5242) 81780390 (5243, 5244) Noticel Brotein cim. Const.	Grand Smr. Galles	39306 (52
0 0/1/2		2618 9123			2621 8025 2622 8178		2623 9163

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2624	91639308 (5247, 5248)	2624   91639308 (5247, 5248) Novel Protein sim. GBank gij3880355 emb CAB05299  - (Z82285) predicted using Genefinder (Caenorhabdilis elegans)		UNCLASSIFIED	56181686, 22278996, 22278997, 22278998, 22278999, 264259, 29331822, 29331824, 56182181, 29331825, 60432289, 29331828, 35698052, 29146499, 66712502, 52644045, 265008, 60433356, 33109954, 21906754, 265010, 265011, 265019, 264448,
					264288, 21906765, 21908766, 21905767, 29148629, 35695917, 265021, 265022, 27486265, 18108370, 60431528, 55811576,
					35695655, 56182323, 18108385, 87168518, 22279002, 18108391
	86452068 (5249, 5250)	86452068 (5249, 5250) Novel Protein sim. GBank gil2887429 dbj BAA24857  -  (AB007887) KIAA0427 [Homo sapiens]		UNCLASSIFIED	264091, 264511, 263981
	16533797 (5251, 5252)	16533797 (5251, 5252) Novel Protein sim. GBank gil487418 (L20302) - actin filament protein [Gallus gallus]		struct	265008
2627	87636823 (5253, 5254)	87636823 (5253, 5254) Novel Protein sim. GBank gij88462[pinj A27307 - proline-rich phosphoprotein (gene PRH1, Db allele) - human		UNCLASSIFIED	22278996, 265007, 265009, 264448, 21906767, 265021, 264558, 87168518
2628	94848254 (5255, 5256)	Novel Protein sim, GBank gij3123552jembjCAA18609j -		UNCLASSIFIED	22278997, 22278999, 264259, 60432049,
		(AL022578) dJ393P12.2 (hypothetical Proline-rich protein KIAA0269 LIKE) [Homo sapiens]			29331822, 29331824, 29331825, 29331827, 35696052, 29331828, 264907, 264909,
					265008, 264591, 60433356, 60433438, 265010, 265017, 265018, 264380, 264288
				ı	18108357, 21906765, 21906768, 265022,
	,				65274791, 264638, 18108387, 87168518,
2629	87376490 (5257, 5258)	2629 87376490 (5257, 5258) Novel Protein sim. GBank		synthase	29331825, 29331826, 264102, 265006,
		gil4929595igblAAD34058.1[AF15182 - (AF151821) CGI-63 protein [Homo sapiens]			264766, 35695917, 264691, 33657023, 263972, 18108374, 22279000
2630	79188364 (5259, 5260)	2630 [79188364 (5259, 5260)]		UNCLASSIFIED	264636, 18108385
2631	94845909 (5261, 5262)	Novel Protein sim. GBank gij321605 pir  JQ1161 - Gag	Contains protein domain (PF00098) - dna_ma_bind	dna_ma_bind	52644507, 52645156, 52646365, 52646842,
		protein - Visna virus (strain EV1)	Zinc finger, CCHC class	l I	22278994, 22278995, 35696286, 56994075,
					22276397, 22776393, 204239, 32643000, 29331822, 29331824, 29331825, 29331826,
					29331827, 29331828, 35696052, 33656970,
					264905, 264509, 264907, 264908, 264511, 264412, 264017, 264008, 264010, 62646317
		`			33657084, 52644296, 265010, 87168559.
					265017, 265018, 265019, 264760, 264762,
					284448, 284288, 284369, 284766, 284768,
					32644229, 219U6/64, 219U6/65, 219U6/66, 1
					52644150, 33657023, 52645129, 33657109,
					33657182, 27486261, 27486262, 27486265,
					33657349, 35695763, 35696423, 65274791,
					35695855, 264634, 264637, 52644332, 56182323, 60432313, 264566, 264486
2632	36730414 (5263, 5264)				264685

22278995, 22278997, 22278998, 264259, 29331822, 29331824, 29331825, 29331827, 264508, 265007, 265008, 265009, 265009, 265010, 265011, 87168559, 265018, 265019, 264683, 264288, 264686, 29148629, 33657023, 264683, 33657182, 35695763, 55811578, 264683, 3652488, 87168518, 22279000, 22279002, 264585	29331826, 263972, 264089	22278997, 264259, 29331824, 66714117, 29331827, 29331828, 264907, 33657084, 265017, 265018, 264448, 264288, 21806766, 21906768, 29148629, 19108376, 55811576, 35695855, 87168518, 22278000	264569, 29331822, 29331828, 265006. 60170831, 264681, 264765, 264685, 29148627, 21906769, 29148784, 265022, 60170615, 264635, 18108385, 56526486, 22279002, 264567	264488, 264489, 52644507, 264887, 5264686, 26278994, 22278998, 22278998, 22278998, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 22278999, 29331822, 29331827, 3264509, 264509, 264509, 264509, 264591, 264592, 264391, 264592, 264591, 264592, 264600, 264600, 264602, 264603, 264764, 264767, 264691, 264764, 264697, 21906768, 21906769, 21906768, 21906769, 21906768, 21906769, 21906769, 246637, 264639, 62474620, 27486264, 18108370, 264693, 5264639, 264639, 264639, 264639, 264639, 264639, 264639, 264639, 264567, 264564, 264568, 264566, 264488, 264567, 264564, 264567, 264639, 264489, 264567, 264564, 264567, 264567, 264639, 264567, 264567, 264567, 264569, 264567, 264567, 264567, 264569, 264567, 26467, 26467, 26467, 26467, 26467, 26467, 26467, 26467, 26467, 26467, 26467, 26467, 26467, 26467, 26467, 26467, 26467, 26467, 26467
	UNCLASSIFIED			phosphalase
Contains protein domain (PF00084) - Sushi domain (SCR repeat)			Contains protein domain (PF01546) - Peptidase family M20/M25/M40	Contains protein domain (PF00782) - phosphatase Dual specificity phosphatase. catalytic domain
	87330921 (5267, 5268) Novel Protein sim. GBank gij5441611]emb CAB46854.1  -  (AJ388555) hypothetical protein [Canis familiaris]	86623144 (5269, 5270) Novel Protein sim. GBank gil46806631gblAAD27721.1 AF13294 - (AF132946) CGI-12 protein [Homo sapiens]	87260534 (5271, 5272) Novel Protein sim. GBank gil3879146jemblCAB07846j - (293386) Simitarity to Yeast hypothetical 52.9 KD protein (SW:P43616); cDNA EST EMBL:M89432 comes from this gene; cDNA EST EMBL:D71008 comes from this gene; cDNA EST EMBL:D73578 comes from this gene; cDNA EST EMBL:D73578 comes from this gene; cDNA EST EMBL:D69025 comes from this gene; cDNA	95011299 (5273, 5274) Novel Protein sim. GBank gil4758208 ref NP_004081.1 pDUSP - dual specificity phosphatase 3 (vaccinia virus phosphatase VH1-related)
3 95011617 (5265, 5266) P	2634 87330921 (5267, 5268) N	2635   86623144 (5269, 5270)   9	2636   87260534 (5271, 5272)   1	2837 95011299 (5273, 5274)

60424179, 52644507, 52646842, 18108398, 56182575, 22278995, 22278996, 35696288, 22278997, 22278999, 264258, 60432049, 22378997, 22278999, 264258, 60432049, 29331822, 6043356, 33657402, 55812038, 55811386, 265019, 2642289, 26440429, 56181652, 29148627, 29148629, 5581450, 29148784, 35695917, 265021, 52644150, 33657023, 65274620, 33657109, 35695763, 18108374, 55810764, 35696423, 55811576, 35695855, 60431850, 56182323, 60432113, 264404	264488, 29331825, 35595052, 264508, 264509, 264509, 264512, 33657402, 60433438, 264758, 85558542, 264600, 285020, 265021, 33657109, 264568, 35696423, 264565, 264486	29331822, 29331826, 264906, 33109954, 265017, 265019, 21906768, 35695763, 264636, 264637, 18108387	264828	264259, 29331822, 33657402, 265019, 264369, 264691, 264634, 56526486, 22279002	22278997, 264259, 29146499, 56182435, 264910, 265010, 18108351, 264682, 284688, 284863, 264690, 33657109, 18108370, 26397, 264690, 33657109, 18108370, 26397, 18108374, 264634, 264557, 264558, 18108385, 264482	264107, 264687	265007	
UNCLASSIFIED	synthase	nelicase	UNCLASSIFIED	UNCLASSIFIED	helicase	nucl_recpt	UNCLASSIFIED	UNCLASSIFIED
Contains protein domain (PF00076) - UNCLASSIFIED RM, RBD, or RNP domain)		Contains protein domain (PF00271) - helicase Helicases conserved C-terminal domain			Contains protein domain (PF00270) - helicase DEAD/DEAH box hellcase	Contains protein domain (PF00651) - Inucl_recpt BTB/POZ domain		-
2636   94326733 (5275, 5276) Novel Protein sim. GBank glk929688igblAAD34105.1 AF15186 - (AF151868) CGI-110 protein [Homo saplens]	95381346 (5277, 5278) Novel Protein sim. GBank gil2190007 dbj BAA20355  - (AB004109) phosphatidylserine synthase II [Cricetulus griseus]	87781330 (5279, 5280) Novel Protein sim. GBank gil3158516 (AF067617) - contains similarity to chromo (chromatin organization modifier) domains (Pfam: chromo.hmm. score: 17.76 and 27.94) and to helicases conserved C-terminal domain (Pfam: helicase_C.hmm, score: 67.00) [Caenorhabditis elegans]		Novel Protein sim. GBank unknown [Mus musculus]	.87643961 (5285, 5286) Novel Protein sim. GBank gil4490304 emb CAB38785.1  - (AL035678) putative protein [Arabidopsis thailana]	88177671 (5287, 5288) Novel Protein sim. GBank gij3789797lgbJAAC67502.1  - (AF059569) actin binding protein MAYVEN [Homo sapiens]	(0)	94148542 (5291, 5292) Novel Protein sim. GBank gil1706722 sp P49749 EVX2_MOUSE - HOMEOBOX EVEN SKIPPED HOMOLOG PROTEIN 2 (EVX-2)
94326733 (5275, 5276)			11669834 (5281, 5282)				17277228 (5289, 5290)	
2638	2639	2840	2641	2842	2643	2644	26	2648

2647	2647 91212978 (5293, 5294)			UNCLASSIFIED	56182575, 22278996, 35696286, 22278998,
					264259, 29331822, 36182181, 29331825, 60424269, 60432289, 35696052, 66712502
					264908, 265007, 55812038, 33109954,
					21906754, 33657084, 265019, 264448.
					264288, 56181562, 21908765, 21906766.
					21906768, 21906769, 35695917, 265020,
					265021, 52644150, 264693, 33657109,
					33657349, 60431528, 18108374, 55810764,
					35696423, 56182323, 60432113, 22279002,
2849	2848 87600587 (F205 6206)				264564
7	0 00000 (0500, 0500)				29146498, 56182435, 33109954, 265011,
					264682, 55811957, 35695917, 264690,
36.40					263976, 18108377, 35696423, 60432113
		194 120/03 (328/, 3288) Novel Protein Sim. GBank gij3041852 (AC004539) -		UNCLASSIFIED	56182575, 35696286, 22278998, 29331824,
		nunction; similar			29331826, 60432289, 66712502, 56182435,
		sapiens			60170831, 60432229, 33657402, 33109954,
					21906754, 265017, 264686, 264688,
_					21906765, 21906768, 60170615, 264693,
					263967, 18108370, 263976, 60170394,
					60432113, 22279002, 264563
7650		87297533 (5299, 5300) Novel Protein sim. GBank gij5360271 dbj BAA81908.1	(		264685
2661	10003 1003/ 37/00/08	(Abuzasas) nrre i -3 [Haiocynthia roretzi]			
	(2006, 1300), 3302)	- 1.1987/43 (3301, 3302) Novel Protein Sim. GBank gij4240225[db] BAA74891.1]	Contains protein domain (PF00054) - synthase	synthase	29331824, 29331826, 29331827, 265007,
		(Abuzuara) KiAAuasa protein [Homo sapiens]	Laminin G domain		55812038, 21906754, 18108366, 18108384,
2652	10343125 (5303, 5304)				26.15004, 204307
2653	87798735 (5305 5308)	Now Designation		UNCLASSIFIED	764697
3	(2005, 2003, 2009)	Nover Profess Sim. Gbank gil4493956 emb[CAB11123.2]		UNCLASSIFIED	265018, 18108370, 18108387, 264566
		(299331) predicted using next con; MAL3P6.28	1		
		model crossiem hundhalical protein, 161, 167, 48; Similarity 10			•
_		D.melanogaster S.cerevisiae & S.nombo) Celegans.	•		
2854	95103240 (5307, 5308)				60424179, 65274572, 56182575, 264259.
					56182181, 264908, 56182435, 55811957,
					35695917, 265021, 263976, 55810764,
3556					65274791, 56182323, 83373044, 65274727
		91228018 (3309, 3310) Novel Protein sim. GBank gij3875272 embjCAB02861   -	_	transcriptfactor	56182575, 56181686, 264092, 264259.
		(281051) predicted using Genefinder; similar to Zinc finger,			56182181, 60432289, 264907, 33657402,
		Canca type (Ring linger); cDNA EST yk443h5,3 comes	(Inger)		55812038, 21906754, 87168559, 265017,
		from this gene; cond EST yk443h5.5 comes from this gene			264448, 264369, 264288, 21906765,
		l Caenornabonis elegans			21906766, 21906767, 21906768, 33657109,
					18108370, 264628, 55811576, 264556,
ĺ					264639, 83373044, 56526486, 264404,
<b>3656</b>	84562601 (5311, 5312)	84562601 (5311, 5312) Novel Protein sim. GBank gil3043718IdbilBAA255231.			200432113
		(AB011169) KIAA0597 protein (Homo sapiens)			250502
			**************************************		

2657	52561728 (5313, 5314)	52561728 (5313, 5314) Novel Protein sim. GBank gil5689509idbjlBAA83038.11 -		dna_rna_bind	264693
		(AB029009) KIAA1086 protein [Homo saplens]			707000 0007000
2658	88062454 (5315, 5316)	88062454 (5315, 5316) Novel Protein slm. GBank gij3688089 (AC005757) -	Contains protein domain (PF00560) - nucleaseinhib	nucleaseinhib	35696286, 264259, 29331822, 29331824,
		R32611_1 [Homo saplens]	Leucine Rich Repeat		29331826, 29331828, 265019, 264663,
					21906768, 35695917, 264593, 35695855,
					204037, 07 1003 10, 204400, 204307
2659	87600755 (5317, 5318)	87600755 (5317, 5318) Novel Protein sim. GBank gi[5420387 emb CAB46679.1 -	Contains protein domain (PF01426) - UNCLASSIFIED	UNCLASSIFIED	264909, 264910, 265018, 264369, 264769.
		(AJ243459) proteophosphoglycan [Leishmania major]	BAH domain		Z1905/09, Z04093, Z0397Z, 10100300
2660	91718472 (5319, 5320)	91718472 (5319, 5320) Novel Protein sim. GBank		kinase	264488, 65274572, 33696286, 22276996.
		gil728837 sp P39194 ALU7_HUMAN - !!!! ALU SUBFAMILY	EF hand		22278999, 264259, 29331822, 29331824,
		SO WARNING ENTRY IIII			60432289, 29331826, 35696052, 264908,
					56182435, 265008, 265009, 60433356,
					264594, 265010, 265018, 55811150,
					18108351, 264682, 264684, 264369, 264288.
					264687, 21906765, 29148784, 35695917,
					60170615, 52644150, 33657023, 33657109,
			,		35696423, 35695855, 264556, 60170394,
					18108385 22279000 22279002
				alveografoin	ENA32049 264259 29331824 29331825.
2661	95342817 (5321, 5322)	95342817 (5321, 5322) Novel Protein sim. GBank		man design	10031636 30331637 30331838 38400R
		gil4758048 ref NP_004739.1 pCPR8 - cell cycle progression			29331626, 29331621, 29331626, 204300.
		8 protein			264909, 264593, 33103954, 265010, 265017,
		-			265018, 265019, 264760, 264448, 264369,
_					264288, 21906765, 21906768, 265022,
		,			264691, 33657023, 27486262, 60431528,
				_	18108374 35695855 18108388 264482
	7				DEJEKE DEJEKE DEJEKE DEJABE
<b>7</b> 992	_				204303, 204330, 204330, 204460
2663		87780623 (5325, 5326) Novel Protein sim. GBank gij3874714 jembjCAA91263 j		dehydrogenase	[264906, 264909, 264757, 264758, 204707,
		(Z66494) similar to choline dehydrogenase; cDNA EST			264691, 3365/023, 264636
		yk346d5.5 comes from this gene; cDNA EST yk346d5.3			:
		comes from this gene [Caenorhabditis etegans]			
2664		85518329 (5327, 5328) Novel Protein sim. GBank gil1389670 (U58977) - Notch	Contains protein domain (PF00008) - oncogene	- oncogene	35696286, 264509, 264595, 264288, 254585,
2		Interior of the control of the contr		INCI ASSIFIED	35696286 22278999 29331822 35696052.
2007		Moves Protein Sim. Grank gijado4400jembjCAD45314.11			264906 264907 264909 264510, 264511,
.,					264512, 264593, 60433438, 265019, 264681,
					21906765, 21906766, 21906767, 21906768.
_			_	-	265020 265022 35696423 35695855.
					22279002 264482 264486
9000	_	Marie Contain aim Comt		INCIASSIFIED	29331825, 265007, 264910, 60432229.
7000	-	orozofiz (3331, 3332) regen fragen sint. Gbank			1256110 254288 21006757 264558
		gi 5106956 gb AAD39906.1 AF11361 - (AF113615)  EU1/EU2 domain_containing protein FUOS (Homo sanions)			22279002
200	ALCOS COCS OCCUPATO	The state of the s	Contract natural domain (DE01138)	Circlesee	264907 29331830 264681 264683 264288.
7007	0/422/20 (3333, 3334)	o/422/20 (3555, 3554) Nover Protein sim. Goank		2000	12505015 264812 264556 264557 264558
		gi[2500570]sp[Q17533]RNPH_CAEEL - RIBONUCLEASE	3. exonbonuciease family		330833033, 204032, 204330, 204331, 201331,   1048559 264567
		PH-LIKE PROTEIN 80564.1			204333, 204000, 204000, 204000

				INCLASSIEIED I	56181688 35696786 22278998 22278999.
8	(0555, 5555) 01/01718	ZOOB   BIZID/10 (3339, 3330) NOVER PLOYEIN SIMI, OBBAINA   BIZID/10/10/10/10/10/10/10/10/10/10/10/10/10/	<u>i</u>		56182181, 29331824, 60424269, 29331825,
_					35696052, 29331828, 66712502, 56182435,
			÷.		60433356, 264758, 21906754, 55811386.
					265011, 87168559, 265017, 265019,
					55811150, 264448, 264369, 264288,
					21906765, 21908768, 21906768, 55811957.
					35695917, 265020, 265021, 33657023,
	•				264692, 33657109, 35695763, 60431528,
					18108374, 35696423, 55811576, 264634,
					60431850, 83373044, 18108385, 87168518.
					22279000, 264563, 264564
2669	95415721 (5337, 5338)	95415721 (5337, 5338) Novel Protein sim. GBank gi[2147012 pir  JC4899 - proline			264489, 264689, 21906767, 65274572.
					56182575, 21906768, 29148627, 21906769,
					29148629, 35696286, 35695917, 22278998,
		,			22278998, 265021, 265022, 60170515,
					52644150, 60432049, 264259, 264691,
					33657023, 264692, 29331822, 29331824,
					29331825, 60432289, 33657109, 29331826,
					29331827, 35696052, 29331828, 29146498.
		-			29146499, 264905, 264908, 52644045,
					264909, 56182435, 35696423, 65274791,
					35695855, 265006, 264910, 264635.
					60432229, 264592, 264638, 56182323,
					60433356, 60170394, 60433438, 264559,
					264595, 55812038, 33109954, 87168559,
					60432113, 265019, 264448, 264369, 264684,
					264288
2670	87613234 (5339, 5340	2670 87613234 (5339, 5340) Novel Protein sim. GBank	Contains protein domain (PF00628) -		18108370, 263974
		gi[1723523]sp[Q10362]YDBB_SCHPO - HYPOTHETICAL	PHD-finger		
2674	91214936 (5341 5342	94.9 ND FRO LEIN OZZE 12.110 IN OTINOMICO MILE I		ransport	52646842, 56994075, 264259, 29331822.
					29331824, 29331825, 29331827, 33656970.
		Iong-chain acyl-CoA synthetase homolog 2: VLCS-H2			264509, 265006, 33109954, 21906754.
		[Homo sapiens]			264682, 264288, 265021, 33657023.
					33657109, 33657182, 27486261, 27486262,
					27486265, 18108376, 18108385
2872	87399123 (5343, 5344	2672   87399123 (5343, 5344) Novel Protein sim. GBank	Contains protein domain (PF01344) - UNCLASSIFIED	UNCLASSIFIED	264767
		gil4966346lgblAAD34677.1 AC00634 - (AC006341)	Kelch motif		
		Contains two PF 01344 Kelch motif domains. [Arabidopsis	•		
		[thaliana]			

264488, 22278996, 35696286, 264091, 264259, 29331824, 29331826, 35696052, 264511, 55812038, 85658542, 264766, 21906765, 35695917, 264629, 35696423, 18108383, 87168518	264691 264006 265008	18108394, 65274572, 56182575, 22278994, 22278995, 56994075, 22278996, 35696286, 22278995, 56994075, 22278996, 35696286, 22278995, 22278996, 2693080, 22278996, 264400, 224259, 22278995, 264400, 224259, 265608, 26509, 26331826, 265009, 60432229, 265007, 265008, 265009, 60433229, 265011, 87168559, 265017, 265019, 265011, 87168559, 265017, 265019, 21906765, 21906768, 21906765, 21906769, 256021, 265020, 265021, 263967, 33657109, 27486264, 27486265, 2365367, 33657109, 27486264, 27486265, 2365367, 33657109, 27486264, 27486265, 2365367, 33657109, 27486264, 27486265, 2365369, 33657109, 22279000, 22279002, 264564
transcriptfactor	UNCLASSIFIED	synthase
Contains protein domain (PF00170) - bZIP transcription factor		Contains protein domain (PF00501) - synthase AMP-binding enzyme
Novel Protein sim. GBank gil4758824 ref NP_004280.1 pNRF3 - nuclear factor (erythroid-derived 2)-like 3		94329600 (5353, 5354) Novel Protein sim. GBank gij 1079042 pir  SS2154 - acetyl-
94847721 (5347, 5348)		2677 94329800 (5353, 5354)
	Contains protein domain (PF00170) - transcriptfactor bZIP transcriptlon factor	94847721 (5347, 5348) Novel Protein sim. GBank gil4758824[ref[NP_004280.1]pNRF3 - nuclear factor gil4758824[ref[NP_004280.1]pNRF3 - nuclear factor (erythroid-derived 2)-like 3 79563835 (5349, 5350) UNCLASSIFIED

264488, 263994, 264489, 18108394, 52646842, 35696286, 22278999, 264259, 29331825, 35696286, 22278999, 264269, 29331825, 35696052, 264508, 264905, 264509, 264908, 264509, 264509, 264909, 26450, 264909, 264510, 265008, 264909, 264510, 265008, 264909, 264500, 264910, 265009, 264601, 265001, 87168559, 265019, 264601, 264602, 264603, 265017, 265019, 264605, 264602, 264603, 265017, 265019, 264605, 264691, 3369717, 264693, 19108364, 18108305, 33657109, 33657349, 264636, 264631, 264632, 264636, 264631, 264632, 264636, 264636, 264636, 264636, 264636, 264636, 264656, 264663, 264656, 264656, 264663, 264566, 26466, 26466, 26466, 26466, 26466, 26466, 26466, 26466, 26466, 264	264488, 22278996, 35696286, 264259, 29331826, 29331826, 264259, 56182435, 264113, 264511, 265008, 6043348, 265611, 265008, 21906766, 21906766, 21906766, 21906769, 25610764, 265021, 264690, 33657023, 55810764, 35696423, 55811576, 264631, 18108381, 60170394, 83373044, 87168518, 264566	264259, 264102, 264905, 264908, 285007, 265008, 33109954, 265011, 18108351, 264768, 33657023, 20281149, 263972, 264630, 264638, 264638	264489, 22278995, 264509, 264805, 264906, 264907, 264907, 264909, 264510, 265006, 264512, 264910, 264594, 60433438, 264758, 264603, 264604, 265019, 264605, 264760, 264764, 266687, 264768, 264769, 21908769, 35695917, 265020, 33657023, 264631, 264638, 264639, 264638, 264686, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264686, 264688, 264638, 264638, 264638, 264638, 264638, 264638, 264688, 264638, 264638, 264638, 264638, 264638, 264638, 264638, 264688, 26488, 26468	35696286, 264908, 55811386, 265017, 55811150, 55811957, 35695917, 60431528, 55810764, 55811576, 35695423, 65274791, 56526486	264909, 264769, 264635, 264636
UNCLASSIFIED	reductase	struct	kinase	- UNCLASSIFIED	- interfeukinrecept
	Contains protein domain (PF00970) - reductase FAD/NAD-binding Cytochrome reductase	Contains protein domain (PF00036) - struct EF hand		Contains protein domain (PF00076) - RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)	Contains protein domain (PF00622) - interleukinrecept SPRY domain
95001694 (5355, 5356) Novel Protein sim. GBank gil86760 pir  A40465 - alanine transaminase (EC 2.8.1.2), cytosolic - human		87800356 (5359, 5360) Novel Protein sim. GBank gil4589604 dbj 8AA76824.1  - (AB023197) KIAA0980 protein [Homo sapiens]	90933844 (5361, 5362) Novel Protein sim. GBank gil728837 sp P39194 ALU7_HUMAN - !!!! ALU SUBFAMILY SQ WARNING ENTRY !!!!	94138934 (5363, 5364) Novel Protein sim. GBank gil423468 pir  JQ1974 - HTF9-C protein - mouse	87774405 (5365, 5366) Novel Protein sim. GBank gi[5114351[gb]AAD40286.1 - [AF156271) RING finger protein terf [Homo sapiens]
2678 95001694 (5355, 5356)	2679   95361544 (5357, 5358)	2680   87800356 (5359, 5360)	2681 <u>90933844 (5361, 5362)</u>	2682   94138934 (5363, 5364)	2683 87774405 (5365, 5366

UNCL Contains protein domain (PF01344) - struct Kelch motif
Contains protein domain (PF00515) - Iranscriptfactor TPR Domain
Novel Protein sim. GBank gij3880023jembjCAA97339j - Contains protein domain (PF00560) - (Z73098) Similarity to yeast hypothetical protein (Swiss Prot Leucine Rich Repeat accession number Q09695j; cDNA EST EMBL:D72982 comes from this gene; cDNA EST EMBL:D75947 comes from this gene [Caenorhabditis elegans]
Contains protein domain (PF00620) - struct RhoGAP domain
Contains protein domain (PF00326) - peptidase Prolyl oligopeptidase family

				ED 29331828, 264512, 264555, 264555, 264557, 264558, 264559, 264558, 264559, 264559, 21906765, 21906768, 265021, 264563
UNCLASSIFIED	transcriptfacto	UNCLASSIFIED		UNCLASSIFIED
-	Contains protein domain (PF00412) - transcriptfactor LIM domain containing proteins		. •	Contains protein domain (PF00642) - Zinc finger C-x8-C-x5-C-x3-H type
2697   91638472 (5393, 5394) Novel Protein sim. GBank gij5689473 dbj BAA83020.11- (AB028991) KIAA1068 protein [Homo sapiens]	94325891 (5395, 5396) Novel Protein sim. GBank gil841318 (U22818) - mutant sterol regulatory element binding protein-2 [Cricetulus griseus]	87780650 (5397, 5398) 94139836 (5399, 5400) Novel Protein sim. GBank gil5174395 ref NP_006006.1 pB120 - Brain protein 120	94148584 (5401, 5402) Novel Protein sim. GBank gil1572801 (U70854) - F38A5.1 gene product [Caenorhabditis elegans]	57295368 (5403, 5404) Novel Protein sim. GBank gil2605967 (AF030027) - 24 [Equine herpesvinus 4] 87649514 (5405, 5406) Novel Protein sim. GBank gil5689399[dbj BAA82983.1] - (AB028954) KIAA1031 protein [Homo sapiens]
91638472 (5383, 5384)		87780650 (5397, 5398) 94139836 (5399, 5400)		
2697	2698	2700	2701	2702

264488, 22278995, 22278998, 29331828, 29146499, 264905, 264908, 264907, 52644045, 264511, 33657402, 264600, 264602, 265017, 264605, 264781, 18108351, 264764, 264687, 264769, 265021, 264691, 18108374, 264634, 264693, 18108370,		UNCLASSIFIED 264488, 52646642, 65274572, 22278994, 56994075, 22278997, 264259, 29331824, 29331826, 29331826, 29331826, 29331824, 29331826, 29331826, 29331826, 29331826, 29331826, 29331826, 29331826, 264909, 5264909, 5264909, 5264909, 5264909, 5264909, 265010, 265011, 265017, 265019, 264681, 26	22278996, 22278998, 56182435, 21906754, 87168559, 265017, 264448, 52645129	UNCLASSIFIED 65274572, 264259, 29331822, 29331825, 60432289, 29331828, 29331827, 29331828, 264909, 264510, 265007, 264910, 60433356, 60433438, 33109954, 265010, 265011, 264389, 264288, 264765, 264693, 264565	UNCLASSIFIED 29331822, 18108370, 18108374, 83373044	29331824, 264759, 264693, 18108382, 18108388
2704   87649515 (5407, 5408) Novel Protein sim. GBank gil4335694 gb AAB63294 - (AF008554) Implantation-associated protein [Rattus norvegicus]	2705   87771745 (5409, 5410)			91011351 (5415, 5416) ,		2710   87627979 (5419, 5420) Novel Protein sim. GBank gil4468311 emb CAB37992  - (AL031432) dJ465N24.2.1 (PUTATIVE novel protein) ((soform 1) [Homo sapiens]

264488, 264687, 52645156, 264769, 21906764, 21908765, 21906767, 21906768, 21906769, 55811957, 56994075, 22278997, 22278998, 265020, 265021, 264690, 264259, 264691, 264692, 33657023, 29331822, 29331827, 27488262, 33657349, 264508, 264905, 264907, 60431528, 264908, 264909, 5581076, 3569642, 33657349, 264508, 265007, 264910, 60431830, 60432229, 18108385, 21906754, 33657084, 87168518, 87168474, 87168559, 60432113, 265017, 22279000, 265018, 265019, 22279002, 264760, 55811150, 264681, 18108351, 264565, 264764, 264568, 264288, 264766	264486, 35696286, 22276998, 264259, 29331824, 60432289, 35696052, 264508, 264906, 66712502, 52644045, 265006, 60432229, 33657402, 60433356, 265006, 265019, 18108351, 264681, 264288, 2506769, 21906766, 21906768, 21906768, 21906769, 25811957, 3569517, 265020, 265021, 60170815, 264691, 264692, 33657023, 33657109, 33657108, 33657139, 18108370, 18108374, 35696423, 35695855, 264555, 52644332, 56182323, 87168518, 60432113	66714117, 264906, 264563	264636 264508, 264905, 264907, 264908, 264909, 264910, 264758, 265011, 264760, 264764, 284288, 264766, 264768, 284636	264091, 264259, 29331822, 66714117, 264908, 264369, 264693, 264556, 264563 264593, 264558	264693 29331822, 87168559, 265019, 265021, 52644150, 264691
UNCLASSIFIED	Iransferase	UNCLASSIFIED	UNCLASSIFIED	glycoprotein	UNCLASSIFIED
	Contains protein domain (PF00515) - transferase TPR Domain				
2711   94111920 (5421, 5422) Novel Protein sim. GBank gij3122400 sp O35682 MUG_MOUSE - MYELOID UPREGULATED PROTEIN		86003064 (5425, 5426) Novel Protein sim. GBank gi[2477513 (AC002398) - F25965_3 [Homo sapiens]		88003068 (5431, 5432) Novel Protein sim. GBank gi[2477513 (AC002398) - F25965_3 [Homo sapiens] 80077461 (5433, 5434) Novel Protein sim. GBank gi[3327046 db] BAA31591  - AB014516  KIAA616 protein Immo sapiens]	
11 94111920 (5421, 5		2713   88003064 (5425, 5	2714   13528218 (5427, 5428) 2715   94122454 (5429, 5430)	2716   88003068 (5431, 5 2717   80077461 (5433, 5	2718 79604062 (5435, 5438) 2719 88180423 (5437, 5438)

2720	2720   95086242 (5439, 5440) Novel Protein sim. GBank	Novel Protein slm. GBank gil1335873 (U46690) - ATP-	Contains protein domain (PF00270) - helicase		18108374, 60424179, 264489, 56182435,
					265020, 265022, 265008, 265008, 264092.
					264638, 60432229, 264691, 264692,
					33657023, 264693, 33657402, 83373044,
					29331824, 18108366, 60424269, 29331826,
					18108385, 52645129, 21906/54, 35696052, (
					29331828, 87168474, 264100, 265010,
			-		265011, 265019, 22279002, 264905, 264482,
					264563, 264906, 18108351, 264681,
					18108370, 29331830, 264908, 66712502,
12.5	7				52644045, 264909, 264828, 18108354
7/9	9224222 (2441, 2442)		Contains protein domain (PF01172) -		22278995, 35696286, 264259, 29331822,
		gija829003[gojAAD34092.1[AF15185 - (AF151855) CGI-97	Uncharacterized protein family		29331824, 66714117, 29331826, 264906,
		protein [Homo sapiens]	UPF0023		60433438, 265017, 18108351, 264448,
			•		264288, 264769, 21906766, 265021,
					33657109, 263969, 60431528, 264629,
				_	55811576, 65274791, 35695855, 264631,
	┰				264637, 60170394, 56182323, 22279000
77/7			Contains protein domain (PF00566) - oncogene	oncogene	35696286, 22278999, 21906754, 265017,
		(AC004997) match to ESTs AA667999 (NID:g2626700).	TBC domain		264762, 264288, 21906765, 21906767,
		AA165465 (NID:g1741481), Z45871 (NID:g575105), and			21906768, 35695917, 18108362, 27486262.
		T84026 (NID:g712314); similar to various tre-like proteins			35695855, 264558, 264559
		Including: AF040654 (PID:g2746883), D13644			
27.23	2723 87387732 (5445 5446)	(rio.92104371), ALOZ114			000000000000000000000000000000000000000
77.7	(0++0' 0++0' 30' 100' 10			UNCLASSIFIED	264508, 264509, 264906, 264909, 264910,
		,	ŀ		55812038, 264768, 264687, 264629, 264636,   264486
2724		87639563 (5447, 5448) Novel Protein sim. GBank		ubiquitin	18108396, 22278999, 20281099, 29331824.
		gil4680681[gb]AAD27730.1]AF13295 - (AF132955) CGI-21			29331826, 60432289, 29331828, 60170831.
		protein [Homo sapiens]			60432229, 60433438, 18108351, 264682.
					21906766, 21906767, 21906769, 35695917,
					33657023, 33657109, 18108372, 18108374,
					35695855, 22279000, 22279002
2725		94853991 (5449, 5450) Novel Protein sim. GBank gij3169705 (AC004780) -		UNCLASSIFIED	264488, 52644507, 264259, 29331827,
		F17127_1 [Homo saplens]			21906754, 285011, 18108351, 264448.
_					264288, 264685, 264689, 35695917, 265020,
					33657182, 27486261, 18108370, 18108374,
					35696423, 18108385, 22279000
2726		86880599 (5451, 5452) Novel Protein sim. GBank gij3342738 (AC005328) - R26660 1, partial CDS (Homo sapiens)		MHC	264488, 264828, 264685

WO 00/58473 PCT/US00/08621

22	91010470 (5453, 5454)	2727   91010470 (5453, 6454) Novel Protein sim. GBank gil7312671splP39219IRLUA_ECOLI - RIBOSOMAL LARGE SUBUNIT PSEUDOURIDNE SYNTHASE A (PSEUDOURIDYLATE SYNTHASE) (URACIL HYDROLYASE)	Contains protein domain (PF00849) - synthase RNA pseudouridylate synthase	synthase	65274572, 56182575, 22278994, 56994075, 22278998, 22278997, 22278999, 60432049, 22331822, 22331822, 22331822, 22331822, 22331827, 35696052, 22331828, 3365909, 66412502, 264910, 33657402, 60433438, 264758, 55812038, 21906754, 23657084, 55811386, 265018, 265019, 244767, 21906765, 21906767, 21906769, 23657109, 33657109, 33657109, 33657109, 3365739, 55811576, 3569623, 33659655, 264630, 60431850, 264636, 56182323, 87168518, 60432113, 22279000.
2728	94126022 (5455, 5456)	2728 94126022 (5455, 5456) Novel Protein sim. GBank gij3880433 emb CAA91399j - (Z66521) similar to mitochondrial RNA splicing MSR4 like protein; cDNA EST EMBL:C09217 comes from this gene [Caenorhabdiis elegans]	Contains protein domain (PF00153) - UNCLASSIFIED Mitochondrial carrier proteins	UNCLASSIFIED	264564, 264365 18108394, 56182435, 21906767, 55811957, 35895855, 265021, 264690, 264556, 264259, 264557, 29331822, 264559, 264448, 264288
2729	94126024 (5457, 5458)	2729   94126024 (5457, 5458) Novel Protein sim. GBank gil2408095 emb CAB16300  - (299168) putative RNA splicing protein [Schizosaccharomyces pombe]	Contains protein domain (PF00153) - UNCLASSIFIED Mitochondrial carrier proteins	UNCLASSIFIED	65274572, 284259, 60432289, 66712502, 56182435, 264448, 264288, 264369, 55811957, 265021, 264557, 60432113
2730	94126026 (5459, 5460)	Novel Protein slm. GBank gij3880433 emb CAA91399  - (Z66521) similar to mitochondrial RNA splicing MSR4 like protein; cDNA EST EMBL:C09217 comes from this gene [Caenorhabditis elegans]	Contains protein domain (PF00153) - transport Mitochondrial carrier proteins	Uansport	264887, 29331824, 29331826, 35696052, 264107, 56182435, 265008, 265009, 264592, 60431735, 265011, 264601, 265017, 18108331, 264288, 29148627, 55811957, 265021, 264690, 18108368, 18108374, 264557, 264558, 264559, 18108387,
2731	87723022 (5461, 5462)	87723022 (5461, 5462) Novel Protein sim. GBank gi 1723239 sp Q10166 YAUB_SCHPO - HYPOTHETICAL 35.7 KD PROTEIN C26A3.11 IN CHROMOSOME I	Contains protein domain (PF00795) - Carbon-nitrogen hydrolase		264259, 35696052, 265006, 264759, 264762, 26448, 264288, 29148627, 21906769, 87168518, 22279002
2732	94126028 (5463, 5464)	94126028 (5463, 5464) Novel Protein sim. GBank gil3880433 emb CAA91399  - (Z66521) similar to mitochondrial RNA splicing MSR4 like protein; cDNA EST EMBL:C09217 comes from this gene [Caenorhabditis elegans]	Contains protein domain (PF00153) - UNCLASSIFIED Mitochondrial carrier proteins	UNCLASSIFIED	264687, 264489, 18108358, 56182435, 264689, 35696423, 55811957, 265021, 265006, 265008, 264910, 265009, 264680, 264555, 264557, 264558, 264559, 18108383, 33657109, 87168518, 265010, 264601, 60432113, 265017, 264905, 24448
2733	97363060 (5465, 5466) 94140286 (5467, 5468)	97363060 (5465, 5466) 94140286 (5467, 5468) Novel Protein sim. GBank gil4519621 dbj BAA75670.1  - (AB017614) OASIS protein [Mus muscufus]	UNCLASSIFIEI Contains protein domain (PF00170) - transcriptfactor bZIP transcription factor	UNCLASSIFIED transcriptfactor	29331825, 254509, 264809 60424179, 52644507, 56182575, 264259. 29331828, 2264907, 264510, 264910, 60433356, 265019, 55811150, 284881, 264763, 264687, 33657182, 18108370, 60431528, 60431850, 56182323, 83373044

-			-4	The second secon
(2)	gb[T21276, gb[T45403, and gb]A4586113 come from this		grycoprotein	zzz/6996, 6043zz89, z93318z/, z9146498, 264108, 264909, 264112, 33657402,
	gene. [Arabidopsis thaliana]			87168474, 265017, 264762, 264448, 264764,
				264684, 21906765, 264693, 33657109,
_				263976, 284638, 264638, 264557, 22279000,
80247655 (5471, 5472)			INCIASSIEIE	222/9002, 26456/ 264006, 264628, 264620, 263078, 264632
				264564
87604526 (5473, 5474)				264690
. 5478)	85731808 (5475, 5476) Novel Protein sim. GBank gi[2558501 dbj BAA22896  -			264488, 265009, 264768, 264691
	(D63850) hepatoma-derived growth factor [Mus musculus]			
5478)	94319834 (5477, 5478) Novel Protein sím. GBank gij5420387jemb CAB46679.1  - [74.124260) proteonbosenbosenbosen i stehmasia maios		UNCLASSIFIED	264684, 83373044, 264566
5480)	04148762 (5470 5480) Naval Drain Praise in Cont 112417261 LEISINIA INGIO			
9	Novel Frotein sim. Gbank gij341/38bjembjCAA/5495j -		UNCLASSIFIED	264488, 56182575, 22278995, 35696286,
	(113197) mid otubule-associated protein, MAP-115 (Mus			22278997, 22278998, 22278999, 264259.
_	musculus)			29331822, 29331824, 29331825, 29331827,
				35696052, 29331828, 29146498, 29331830,
				265006, 265007, 265009, 60432229,
_				33657402, 55812038, 87168474, 265010.
		,	•	265011, 265017, 265018, 265019, 264605.
				264684 264288 264260 62644220
				200001, 204200, 204309, 32644229,
				71906/65, 21906/66, 21906/67, 21906/68,
				21906769, 265020, 265022, 264691, 264692,
				33857109, 18108370, 18108374, 55810764,
				35695855, 264634, 60431850, 264839,
				56182323, 18108382, 18108385, 65274727,
1				22279002, 264564
(7060	0001 310 (3401, 3402) NOVEL Protein Sim. GBank gij3242764 (AC005154) - similar		UNCLASSIFIED	22278998, 52644045, 52644229, 21906768,
	to protein U28928 (PID:g861306) [Homo sapiens]			21906769, 265020, 60170815, 264691
7484)		_	Iransferase	264259, 264905, 264758, 55812038, 264369,
	gil4758412[ref]NP_004472.1[pGALN - UDP-N-acetyl-alpha-			29148627
	D-galactosamine:polypeptide N-	_	_	
	acetylgalactosaminyltrans(erase 2 (GaINAc-T2)			
5486)	Novel Protein sim. GBank gil4468311 emblCAB379921 -		UNCLASSIFIED	35696286 264259 264906 264908 265006
	(AL031432) dJ465N24.2.1 (PUTATIVE novel protein)			60433438 265017 18108351 264448
	(isoform 1) [Homo sapiens]		-	264764, 264288, 21906765, 21906767
				264690, 264691, 264693, 263969, 263971
				SECRET SEARS SEARED ACTORSOS
		. "		50453443
5488)	94126030 (5487, 5488) Novel Protein slm. GBank gil3880433lemblCAA913991.	Contains protein domain (PE00153) - kinasa	kinseo	18108374 264488 K6182426 21006766
•	(266521) similar to mitochondrial RNA splicing MSR4 like	Mitochondrial carden profess	200	15100014, 204406, 30102433, 21300103, 34606423, 34604047, 34604844, 364020
	protein; cDNA EST EMBL:C09217 comes from this gene			265021 264511 265009 264490 264556
	[Caenorhabditis elegans]			264259, 264557, 56182323, 264558, 264559
				18108383, 29331824, 18108385, 33657109
				29331828 21908754 29331827 29331828
				10001020, 11000101, 10001021, 10001020,
				CURACY OFFICE OF COOL OF MANY CONTROL

2746   67740125 (5489, 5490)   Novel Protein sim. GBank   91403759[glyAD19629]   Contains protein domain (PF00220) - UNCLASSIFED	35696286, 264509, 264905, 264907, 264908, 264909, 264510, 264512, 265008, 264758, 264601, 265117, 264604, 264763, 264628, 264686, 26469, 264693, 35696423, 3569585, 264634, 264638, 264563, 264564, 264636, 264563, 264564,	22278996, 22278998, 22278999, 29331822, 29331826, 29331827, 35696052, 29331828, 264905, 264906, 264907, 264908, 264909, 55644045, 265006, 60170831, 264598, 55812038, 265018, 264683, 264288, 21906765, 21906765, 21906769, 33657102, 264693, 33657109, 18108368, 18108374, 264558, 18108368, 284563	264569, 52644507, 18108394, 22278995, 35696286, 22278997, 22278999, 52645080, 29331824, 56182181, 29331826, 29331827, 35696052, 264907, 264908, 264909, 265009, 33109954, 55811386, 87168474, 265010, 87168559, 264603, 265019, 264760, 264688, 264762, 33657023, 52645129, 27486264, 60431528, 18108374, 35696423, 35695855, 264556, 5618232, 38108385, 264482	65274572, 29331828, 264112, 264511, 265019, 264760, 264767, 264768, 264769, 21908768, 21906769, 265020, 27486262, 56526488, 87168518, 22279000	264259, 264908, 264510, 265008, 265009, 264760, 264369, 264768, 264563	264488, 52644507, 18108398, 56994075, 264259, 29331825, 29331826, 29331827, 29331828, 264508, 265009, 264910, 264591, 264595, 33657084, 265011, 265019, 18108351, 264288, 264686, 264769, 26481957, 264693, 27486264, 18108370, 264563	29331822, 29331824, 265017, 33657023
	helicase	UNCLASSIFIED	glycoprotein			UNCLASSIFIED	UNCLASSIFIED homeobox
489, 5490) Novel Protein sim. GBank gil4405795 gbJAAD19826  - (AF038963) RNA helicase [Homo saplens]  491, 5492) Novel Protein sim. GBank gil4758738 raf NP_004680.1 pMTA1 - metastasis associated 1  493, 5494) Novel Protein sim. GBank gil4191272 emb CAA09984  - (A_012295) apaG protein [Rhizobium etti]  495, 5496) Novel Protein sim. GBank gil4191272 emb CAA09984  - (A_012295) apaG protein [Rhizobium etti]  497, 5498)  499, 5500) Novel Protein sim. GBank gil1872498 (U74297) - PiUS    60ryctolagus cuniculus   60ryctolagus cuniculus   F501, 5502    F503, 5504) Novel Protein sim. GBank gil3041859 (AC004534) - OG-2 homaodomain protein-like; similar to U85087 (PID:91575526) [Homo sapiens]	Contains protein domain (PF00271) - Helicases conserved C-terminal domain	Contains protein domain (PF00320) - GATA zinc finger		Contains protein domain (PF00546) - F-box domain.	,		Contains protein domain (PF00046) Homeobox domain
10 IIO IKI IKI IKI IKI IKI IKI IKI		l - :	اما		5497, 5498)	5499, 5500) [Novel Protein sim. GBank gij1872498 (U74297) - PiUS [Orydolagus cuniculus]	5501, 5502) 5503, 5504) Novel Protein sim. GBank gij3041859 (AC004534) - OG-2 homeodomain protein-like; similar to U65067 (PID:g1575526) [Homo sapiens]

56182575, 56994075, 22278998, 29331822,	29331824, 29331825, 29331826, 265007, 264593, 55812038, 33109954, 18108351, 264288, 56181562, 21906767, 21906768, 255021, 264693, 18108374, 65274791, 264632, 56182323, 22279002, 264583, 264567,	22278999, 66714117, 29331827, 35696052, 29331828, 264906, 264909, 265011, 265017, 265018, 265019, 264288, 21906765, 21906767, 265022, 33657023, 284693, 26182323, 18108382, 22278000		264594	85658542, 264693	264693	65274572, 35696286, 66714117, 29331828, 264508, 56182435, 21906754, 55811957, 264629, 264636, 56182323, 22279002	264908	22278998, 29331822, 29331830, 265010, 265019, 265019, 264288, 21906765, 21906768, 21906769, 265020, 56182323, 22279002, 264563	18108394, 22278998, 264906, 264909, 265006, 2650011, 265006, 265010, 265011, 265017, 265019, 18108351, 264448, 284683, 264686, 264768, 265020, 265021, 265022, 264691, 18108362, 264693, 18108362, 264693, 18108368, 18108370, 18108381, 18108382, 18108382, 18108382, 18108382, 18108382, 18108388, \$7168518	264369, 35696423	52646842, 264259, 29331822, 29331825, 29331826, 29331828, 33656970, 264905, 264907, 29331830, 265006, 265009, 21906754, 265019, 264448, 21906769, 27486262, 56182323, 56526486, 87168518, 264487
UNCLASSIFIED	1	struct	transcriptfactor	UNCLASSIFIED	de -	struct		UNCLASSIFIED	nudease	struct	transcriptfactor	
Contains protein domain (PF00646) -	F-box domain.	aj[535428 (U13736) - calmodulin-   Contains protein domain (PF00036) -   struct     EF hand	Contains protein domain (PF00096) - transcriptfactor Zinc finger, C2H2 type		Contains protein domain (PF00017) - eph Src homology domain 2				Contains protein domain (PF00023) - nuclease Ank repeat	Contains protein domain (PF00989) - struct PAS domain	Contains protein domain (PF01352) - transcriptfactor KRAB box	Contains protein domain (PF00370) - FGGY family of carbohydrate kinases
2753   94138972 (5505, 5506)   Novel Protein sim. GBank gil3851648 (AF098301) - neural F Contains protein domain (PF00646) -   UNCLASSIFIED	box protein NFB42 [Rattus norvegicus]	94115513 (5507, 5508) Novel Protein sim. GBank gi[535428 (U13736) - calmodulin- like protein [Pisum sativum]	5510) Novel Protein sim. GBank gi[2996653 (AC004510) - R30385 2 [Homo sapiens]		Novel Protein sim. GBank src-like adapter protein; nr protein containing SH3 an mouse SLAP; Method: cor author [Homo saplens]	78637846 (5515, 5516) Novel Protein sim. GBank gij3608372 (AF053768) - brain specific cortactin-binding protein CBP90 [Rattus norvegicus]	91005312 (5517, 5518) Novel Protein sim. GBank gi 2072200 (U94863) - p40 [Boma disease virus]	55201	2761 87839597 (5521, 5522) Novel Protein sim. GBank gil4914573 emb CAB43685.1  - (AL050390) hypothetical protein [Homo sapiens]	67592689 (5523, 5524) Novel Protein sim. GBank gij3136150 (AF050182) - PERIOD 3 [Mus musculus]	87539968 (5525, 5526) Novel Protein sim. GBank gi[3511122 (AF060503) - zinc finger protein Homo sapiens	94305140 (5527, 5528) Novel Protein sim. GBank gi[2905843 (AF045244) - nbitol kinase [Klebstella pneumoniae]
53  94138972 (5505, 55		2754 94115513 (5507, 56	2755 88001472 (5509, 5510)	756 11465908 (5511, 5512)	2757 95381590 (5513, 55	2758 79637846 (5515, 50	2759 81005312 (5517, 5	780 79824798 (5519 S	761 87639597 (5521, 5	2762 87592699 (5523, 5	2763 87539968 (5525, 5	2764 94305140 (5527, 8

264488, 56182575, 264259, 29331822, 29331824, 66714117, 29331825, 60432289, 29331828, 264908, 29331830, 56182435, 264592, 33657402, 264448, 264369, 264288, 60170615, 264691, 33657023, 264692, 33657109, 18108374, 55811576, 264634, 264638, 56182323, 83373044, 60432113	18108398, 22278996, 22278996, 22278999, 22278996, 22278996, 22578996, 22578996, 22578996, 22578996, 22578996, 22578996, 22578996, 22578996, 22578996, 18108354, 225789388, 18108388, 18108388	264369	244259, 29331827, 29331824, 090114 117, 60432289, 29331827, 264905, 265009, 264592, 55812038, 6527444, 264766, 21906769, 33657109, 263978, 264553, 264564, 264486	18108374, 254685, 254681, 255370, 56182370, 56182423, 254689, 55810764, 21905766, 358195423, 254689, 55810764, 21905766, 35811957, 358595855, 264110, 255021, 254111, 255022, 255006, 255009, 264092, 264092, 264092, 264691, 264597, 264691, 264592, 263981, 264597, 56182181, 55812038, 29331825, 60424269, 18108385, 29331826, 254107, 60432113, 265017, 55811386, 264107, 60432113, 265017, 55811150, 18108351, 264681, 264906, 18108370, 264484, 264682, 20281069, 264288, 264684, 264683, 264764, 264288, 264684, 264683, 264764, 264288, 264684, 264683, 264764, 264288, 264684, 264568, 264764, 264288, 264684, 264768, 264388, 264764, 264288, 264684, 264768, 264388, 264768, 264388, 264388, 264868, 264388	22278995, 35695486, 22278995, 22278995, 22278995, 2264259, 29331826, 60432289, 35696052, 264112, 33657402, 21906754, 87168559, 265017, 265018, 264289, 21906766, 21906767, 21906769, 356958917, 265020, 2653976, 35696423, 35695855, 87168518, 22279000, 264482
tubulin	kinase	UNCLASSIFIED	histone	UNCLASSIFIED	- ribosomalprot
	Contains protein domain (PF00400) - kinase WD domain, G-beta repeat		Contains protein domain (PF00650) - Histone deacetylase family		Contains protein domain (PF00380) - inbosomalprot Ribosomal protein S9/S16
2769   95311088 (5537, 5538) Novel Protein sim. GBank gil5419859 emb CAB46375.1  - (AL096725) hypothetical protein [Homo sapiens]	17.1  - es	88084071 (5541, 5542) Novel Protein sim. GBank gil3093433 (AC004125) - Unknown gene product [Homo saplens]	95357308 (5543, 5544) Novel Protein sim. GBank gij4885531 ref NP_005465.1 pNY Contains protein domain (PF00850) - histone C - histone deacetylase 5 Histone deacetylase family	94138994 (5545, 5546) Novel Protein sim. GBank gil3288888 (AC005253) - R26445_1 [Homo sapiens]	87819908 (5547, 5548) Noval Protein sim. GBank gil465852 sp P34388 YLS3_CAEEL • HYPOTHETICAL 70.7 Ribosomal protein S9/S16 KD PROTEIN F09G8.3 IN CHROMOSOME III
95311088 (5537, 5538) N	87730182 (5539, 5540)				
2769	2770	2771	2772	2773	2774

264488, 22278995, 56994075, 22278996, 35698286, 22278998, 22278999, 264259, 29331822, 29331824, 29331825, 60432289, 29331827, 29331827, 29331827, 29331827, 29331827, 29331827, 255019, 264448, 264682, 264689, 264689, 266488, 18108354, 52644229, 264689, 21906765, 21906766, 21906767, 255021, 265022, 60170615, 264691, 18108370, 35696423, 56274791, 35695855, 264634, 60431850, 60170394, 56182323, 264558, 264488, 264567	56182575, 22278998, 22278999, 264259, 29331822, 29331824, 264908, 29331830, 264510, 33657402, 21906754, 55811386, 265017, 265019, 284448, 224288, 21906765, 21906766, 21906764, 21906768, 1806768, 1806502, 265020, 265021, 265022, 60170615, 52779000		ED 264907, 264593, 264760, 264628		29146499, 65274791, 264634, 264639	60424179, 29331824, 60424269, 66714117, 29331828, 56182435, 87168474, 265017, 264764, 56181562, 21906765, 21906766, 21906768, 35695917, 265020, 33657023, 35695855, 56182323, 87168518	65274572, 22278996, 22278998, 60432049, 264259, 29331822, 29331824, 29331826, 264259, 29331822, 29331824, 29331826, 60432289, 29331822, 29331824, 29331826, 60432289, 29331822, 264907, 264908, 264909, 265018, 264691, 265019, 264691, 265021, 264691, 18108362, 264692, 2365721, 264628, 263978, 35693685, 264528, 264628, 28657323, 264628, 263978, 35693855, 264527, 264628, 23367109, 33657399, 18108370, 264628, 263978, 35693855, 264527, 264528, 23367109, 33657398, 264528, 23367109, 33657398, 264528, 263978, 36593855, 264527, 264628, 263978, 36593855, 264528, 263978, 36593855, 264528, 263978, 36593855, 264528, 263978, 36593855, 264528, 263978, 36593855, 264528, 263978, 36593855, 264528, 263978, 3659385
ribosomalprol		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	kinase	ubiquitin	struct
Contains protein domain (PF00177) - ribosomalprot Ribosomal protein S7p/S5e				·	Contains protein domain (PF00780) - kinase CNH domain		Contains protein domain (PF00560) - struct Leucine Rich Repeat
2775 95307987 (5549, 5550) Novel Protein sim. GBank gl 4689132 gb AAD27775.1 AF07704 - (AF077042) 30S ribosomal protein S7 homolog [Homo sapiens]	1552)	1554)	1556)	87649729 (5557, 5558) Novel Protein sim. GBank gil4680711[gb]AAD27745.1[AF13297 - (AF132970) CGI-36 protein [Homo sapiens]	94679397 (5559, 5560) Novel Protein sim. GBank 94679397 (5559, 5560) Novel Protein Sim. GBank	5562) Novel Protein sim. GBank gil4469352 gbJAAD21222  - (AF069502) ubiquilin specific protease UBP43 [Mus musculus]	94233146 (5563, 5564) Novel Protein sim. GBank gil4505013 ref NP_002310.1 pLRN  - leucine-rich neuronal protein
95307997 (5549, 5:		79818729 (5553, 5554)			94679397 (5559, 5:	91220057 (5561, 5;	94233146 (5563, 5;
2775	2776	2777	2778	2779	2780	2781	2782

2783	BOOTERSO (REEK EERS)	2783 IBODIEGO VEREE RECEITION Developation Connection			100000 000000 000000
		gij728831 spip39188 aLU1_HUMAN - IIII ALU SUBFAMILY J WARNING ENTRY IIII			zułaus, kułuka, kusara, kasso i
2784			•		264259, 29331822, 29331824, 29331825, 264482
2785		88071930 (5569, 5570) Novel Protein sim. GBank gi 2134933 pir  S58890 - collapsin response mediator protein - human		ASSIFIED	22278999, 264908, 264769, 265018, 264769, 21906765, 21906768, 21906769, 265020, 264564
2786	87408542 (5571, 5572)		gi 2073564 (U80223) - eukaryotic Contains protein domain (PF00069) - kinase kinase; DGCN2 [Drosophila Eukaryotic protein kinase domain		264905, 265017
2787	87901266 (5573, 5574)	87901266 (5573, 5574) Novel Protein sim. GBank gil5174507 ref NP_006020.1 pMA1  - paraneoplastic neuronal antigen		UNCLASSIFIED	264768, 21906765, 21906768, 55811957, 22278999, 264093, 60170615, 264259, 29331822, 18108365, 29331824, 33657109, 29331827, 35696052, 264100, 264105, 264908, 263977, 55811576, 264635, 264637, 60433438, 60432113, 265017, 265019, 22279002, 55811150, 264369, 264288
2788	88090644 (5575, 5576)	88090644 (5575, 5576) Novel Protein sim. GBank gij3252826 (AC004382) - Unknown gene product [Homo sapiens]		UNCLASSIFIED	
2789	85491275 (5577, 5578)	85491275 (5577, 5578) Novel Protein sim. GBank gi Z495729 sp Q92556 Y281_HUMAN - HYPOTHETICAL PROTEIN KIAA0281 (HA6725)		UNCLASSIFIED	264103, 21906769, 264693
	87602784 (5579, 5580)	Novel Protein sim. GBank gi 5101772 emb CAB45135.1  - (AJ242978) p621 [Homo sapiens]		stuct	264488, 264769, 264910, 264631, 264636, 264690, 264691, 264259, 264638, 29331824, 264508, 284509, 264905, 264563, 264906, 264628, 18108370, 264907, 264764, 264908, 264288, 284909
2791	88083195 (5581, 5582)	88083195 (5581, 5582) Novel Protein sim. GBank gi[2911266 (AC002550) - Unknown gene product (Homo sapiens)			21906764, 18108368
	95083783 (5583, 5584)	Novel Protein sim. GBank gi[2854163]gb[AAC02581.1] - (AF045642) No definition line found [Caenomabditis elegans]		,	22276996, 22276997, 264259, 29331822, 29331824, 29331825, 66714117, 29331826, 60432289, 29331827, 35696052, 264906, 66712502, 29331830, 264909, 6043229, 60433356, 60433438, 33109954, 255010, 255011, 255017, 265018, 265019, 26488, 26448, 284289, 264369, 264509, 265021, 3960576, 21906766, 255020, 255021, 35695855, 264636, 264558, 264558, 264558, 264558, 264558, 5662523, 33333044, 18108385, 56526486, 6043313, 22279000, 264567
2793	87425476 (5585, 5586)			UNCLASSIFIED	264259, 60432289, 66712502, 265009, 284836
2784	85794830 (5587, 5588)			UNCLASSIFIED	264689, 265022, 264691, 18108388, 264567

62274572, 56182575, 35696286, 22278998, 22278999, 60432049, 264259, 29331822, 29331822, 29331822, 29331822, 29331827, 36596052, 29331828, 29331827, 36596052, 29331828, 264312, 265007, 264512, 264907, 60432289, 60433356, 60433438, 33109954, 85658542, 265018, 265019, 264288, 264686, 21906764, 21906765, 21906768, 21906769, 25641957, 265020, 265021, 265022, 32644150, 33657023, 264693, 65274620, 33657109, 3569423, 55811576, 6527491, 56182323, 5652686, 60432113, 22279002, 264482, 264593, 264694, 264567, 264482,	181108334, 62474574, 30 162370, 30 52370, 37 12278999, 264490, 60432049, 264259, 29331822, 29331824, 29331824, 35695652, 264509, 264909, 264907, 264909, 66712502, 56182435, 264510, 265009, 264510, 265007, 265008, 264512, 265007, 265008, 264510, 265007, 265009, 264510, 265007, 265001, 265017, 265018, 265019, 265010, 265011, 265017, 265018, 264689, 264682, 264682, 21906766, 21906767, 21906768, 21906766, 21906767, 21906768, 21906766, 21906767, 265020, 265021, 60170615, 2644160, 265021, 60170615, 2644160, 265021, 60170615, 2644160, 265021, 60170615, 2644160, 265020, 265021, 60170615, 2644160, 264562, 264562, 33657182, 34657182, 34657182, 3465724, 35696423, 65274721, 87168518, 22278000, 264563, 264564, 264565, 264567, 264567, 264565, 264567, 264567, 264565, 264567, 264565, 264567, 264567, 264565, 264566, 264567, 264565, 264566, 264567, 264565, 264566, 264567, 264565, 264566, 264567, 264566, 264567, 264565, 264566, 264567, 264565, 264567, 264565, 264567, 264687, 264567, 264687, 264567, 264567, 264687, 264567, 264687, 264567, 264687, 264567, 264687, 264567, 264687, 264687, 264687, 264687, 264687, 264687, 264	56182575, 22278995, 22278996, 22278997, 29331827, 29146499, 264509, 264906, 26182435, 2644757, 21906754, 265010, 265017, 265019, 264681, 264682, 21906765, 21906767, 21906787, 21906789, 21906789, 284690, 33657182, 264629, 18108376, 56182323, 22278002, 264563	264488, 264490, 264259, 264448, 20281149, 20281152, 264556, 264557, 264558, 264559, 264483, 264488, 264567
	UNCLASSIFIED	- UNCLASSIFIED	UNCLASSIFIED
Contains protein domain (PF00179) - ubiquitin Ublquitin-conjugating enzyme		Contains protein domain (PF00569) - UNCLASSIFIED Zinc finger present in dystrophin, CBP/p300	
8. 1pUBE2 - ubiquitin-conjugating us to yeast UBC4/5)	94848857 (5591, 5592) Novel Protein sim. GBank gl 4680651 gb AAD27715.1 AF13294 - (AF132940) CG1-06 protein [Homo sapiens]	95110790 (5593, 5594) Novel Protein sim. GBank gild838557igb AAD31040.1  - (AF143859) potasslum channel modulatory factor DEBT-91 [Mus musculus]	86198005 (5595, 5596) Novel Protein slm. GBank gi 2852645 (AF007160) - unknown [Homo sapiens]
2795   95334868 (5589, 5590) Novel Protein sim. GBank gij5454146jreflNP_006346 enzyme E2E 3 (homologo	2786 94848857 (5591, 5592) P	2797 95110780 (5593, 5584)	2788 86198005 (5595, 5586)

22278996, 22278997, 22278998, 22278999, 60432049, 264259, 29331826, 35696052, 29331828, 264489, 6043356, 33657402, 33109954, 87168474, 264448, 52644229, 21906766, 21906767, 21906768, 35695917, 265020, 265021, 265022, 52644150, 35695855, 264634, 60432113, 22279000	264488, 56994075, 2644259, 20281089, 29331825, 29331827, 264905, 56182435, 265006, 265011, 87168559, 265017, 265019, 264488, 264286, 27486265, 264686, 60170615, 264636, 264557, 264558, 264559, 87168518, 264564, 264566, 264567	265007, 264687	264448, 35695855	264639	264566	264092, 264259, 293318224, 29331827, 29331828, 264508, 265007, 265009, 21908754, 264369, 264288, 264632, 60170394, 264563, 264482	22278995, 22278996, 22278999, 264259, 29331822, 29331824, 29331826, 29331826, 29331827, 29146498, 265008, 265009, 60433438, 265017, 265018, 265019, 264448, 264288, 21906765, 21906767, 21906769, 29148629, 265022, 52644150, 56182323
UNCLASSIFIED	glycoprotein		transport	UNCLASSIFIED	peptidase		UNCLASSIFIED
		Contains protein domain (PF00627) - UBA domain	gi[2337865 (AC002464) - organic   Contains protein domain (PF00083) - Itransport milarity to JC4884 (PID:g2143892)   Sugar (and other) transporter		Contains protein domain (PF01585) - peptidase G-patch domain		*
2799   88090651 (5597, 5598) Novel Protein sim. GBank gij3252825 (AC004382) Unknown gene product [Homo sapiens]	2800   88316481 (5599, 5600) Novel Protein sim. GBank gil4240301 dbj BAA74929.1  - (AB020713) KIAA0906 protein [Homo saplens]		88082477 (5603, 5604) Novel Protein sim. GBank gi[2337865 (AC002464) - organic   Contains protein domain (PFC cation transporter, 50% similarity to JC4884 (PID:g2143892) Sugar (and other) transporter [Homo sapiens]		2804 [57111131 (5607, 5608) Novel Protein sim. GBank gil4559368[gb]AAD23029.1AC00658 - (AC006585) hypothetical protein [Arabidopsis thaliana]		2806   87898951 (5611, 5612) Novel Protein sim. GBank gi 1168973 sp P44403 CLPB_HAEIN - CLPB PROTEIN
88090651 (5597, 5598)	88316481 (5599, 5600)	2801 86068814 (5601, 5602)	88082477 (5603, 5604)	79577446 (5605, 5606)	57111131 (5607, 5608)	87398486 (5609, 5610)	87898951 (5611, 5612) ,
2799	2800	2801	2802	2803	2804	2805	2806

UNCLASSIFIED 52644507, 52646842, 18108398, 56182575, 22278994, 22278994, 22278995, 22278995, 22278996, 26934075, 35696286, 22278997, 22278999, 22278999, 264259, 52845080, 29331822, 29331824, 29331825, 29331822, 29331828, 33656970, 264100, 264105, 264907, 52644045, 6043336, 264506, 28108954, 33657084, 52646317, 21808754, 33109954, 33657084, 52646317, 21808754, 33109954, 265412, 265011, 87188559, 265017, 265018, 265011, 2644229, 21906765, 21906767, 21906768, 21906765, 21906765, 21906767, 265021, 265022, 52644150, 33657023, 52645129, 33657102, 33657182, 27486262, 27486265, 35695855, 52644332, 18108377, 356996423, 35695855, 52644332, 833373044, 18108385, 18108387, 87168518, 8716851	MHC 60432289, 264510, 265010, 265017, 265018, 264686, 265021, 264690, 22279000, 264566	ATPase_associated 18108351	UNCLASSIFIED 284569, 22278996, 264091, 264259, 29331828, 29146499, 29148629, 29148784, 284693	UNCLASSIFIED [22278999, 264259, 66712502, 264693	UNCLASSIFIED 264106	UNCLASSIFIED 56182575, 29331828, 264906, 66712502, 55811388, 265017, 265018, 265019, 264683, 265020, 87168518, 60432113		UNCLASSIFIED   264907 264909	
liar to		nilar 3098			aich .	<u>.</u>	Contains protein domain (PF00170) - transcriptfactor		÷
91720702 (5613, 5614) Novel Protein sim. GBank gil4488310 emb CAB37991  - (AL031432) dJ465N24,1 (PUTATIVE novel protein similar to predicted yeast and worm proteins) [Homo sapiens]	95359111 (5615, 5616) Novel Protein sim. GBank gil5541863 emb CAB51071.1  - (AL096857) hypothetical protein [Homo sapiens]	88083530 (5617, 5618) Novel Protein sim. GBank gi[2772561 (AC004002) - similar to ciliary dynein beta heavy chain; 78% Similarity to P23098 (PID:g118965) [Homo sapiens]			88093334 (5623, 5624) Novel Protein sim. GBank gij3264583 (AC005189) - match to ESTs H97758 (NID:g1118643) and AA085546 (NID:g1628773) [Homo sapiens]	91218755 (5625, 5626) Novel Protein sim. GBank gil4240273 dbj BAA74915.1  - (AB020699) KIAA0892 protein [Homo saplens]	90980906 (5627, 5628) Novel Protein sim. GBank gij3548791 (AC005620) - R33590 1 [Homo sapiens]		95358229 (5631, 5632) Novel Protein sim. GBank gij5420389jemb CAB46680.1  - (AJ243460) proteophosphoglycan [Leishmania major]
91720702 (5613, 5614)			87259032 (5619, 5620)	_			90980906 (5827, 5628)	79774521 (5629, 5630)	95358229 (5631, 5632)
2807	2808	2809	2810	둟	2812	2813	2814	2815	2816

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	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	complement	UNCLASSIFIED	complement	UNCLASSIFIED		
	Contains protein domain (PF00023) - UNCLASSIFIED Ank repeat		Contains protein domain (PF00919) - UNCLASSIFIED Uncharacterized protein family UPF0004	Contains protein domain (PF00386) - complement C1q domain		Contains protein domain (PF00386) - complement C1q domain			
2817 87749542 (5633, 5634) Novel Protein sim. GBank gil1293846 (U56966) - coded for by C. elegans cDNA yk30b3.5; coded for by C. elegans cDNA yk30b3.3 [Caenorhabditis elegans]	2818   88073579 (5635, 5636) Novel Protein sim. GBank gil549986 (U13149) - possible apospory-associated protein [Pennisetum ciliare]		2820   87765744 (5839, 5640) Novel Protein sim. GBank gil4929773igblAAD34147.1JAF15209 - (AF152097) CGI-05   Uncharacterized protein family protein [Homo sapiens] UPF0004	2821   95320511 (5641, 5642) Novel Protein sim. GBank   Contains pro gij399144 sp P02747 C1QC_HUMAN - COMPLEMENT C1Q C1q domain SUBCOMPONENT, C CHAIN PRECURSOR	94260221 (5643, 5644)	2823 85320513 (5645, 5646) Novel Protein sim. GBank gij399144[sp P02747 C1QC_HUMAN - COMPLEMENT C1Q C1q domain SUBCOMPONENT, C CHAIN PRECURSOR		19742170 (5649, 5650)	2826   94311905 (5651, 5652) Novel Protein sim. GBank gij3659683 emb CAA22020  - (AL033503) conserved hypothetical protein [Candida albicans]

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264488, 263994, 264489, 65274572, 29331822, 66714117, 29331827, 29331828, 264508, 264905, 264509, 264906, 264907, 264908, 264909, 264510, 285006, 264511, 265007, 264512, 265008, 264910, 264591,	60432229, 264592, 264593, 264595, 264596, 21906754, 265011, 264600, 264601, 264602, 265017, 264605, 264268, 264766, 264767, 264608, 264768, 264767, 264690, 264600, 26	264951, 264631, 264633, 2031(2, 204023, 35695855, 264631, 264633, 264635, 244555, 264638, 264637, 264556, 284638, 264558, 3373044, 18108385, 60432113, 22279002, 264563, 264564, 264565, 264566, 264567, 264486, 18108391	29331822, 35696052, 264104, 264908, 265007, 264591, 265010, 265017, 265019, 264766, 264686, 55811957, 18108370, 18108374, 35810764, 35696423, 55811576, 6269333	2278996, 22278997, 264091, 264093, 60432049, 264259, 29331822, 29331825, 29331827, 29331827, 29331827, 29331827, 294909, 264510, 264511, 264509, 60433438, 21906754, 265011, 264603, 18108351, 264288, 21906765, 21906768, 21906769, 20448629, 52644150, 264693,	33657109, 18108374, 264634, 18108385, 60432113, 22279000, 264565, 264486 265008, 265019, 264639, 22279002	265017	22278997, 22278999, 52646311, 264286, 264688, 21906767, 60431528, 264638, 22279000	264555, 264556, 264558	263972	ATPase_associated   60432289, 29331828, 265008, 265010, 265017, 26448, 55811957, 265020, 18108370	264259, 264112, 265010, 264762, 264764, 263974, 264555, 264558, 264559
						UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED		ATPase_associated	transcriptfactor
Contains protein domain (PF00386) - complement C1q domain			Contains protein domain (PF00102) - phosphatase Protein-tyrosine phosphatase								Contains protein domain (PF00412) - transcriptlactor LIM domain containing proteins
2827 85320519 (5653, 5654) Novel Protein sim. GBank gij399144 sp P02747 C1QC_HUMAN - COMPLEMENT C1Q C1q domain SUBCOMPONENT, C CHAIN PRECURSOR			91229615 (5655, 5656) Novel Protein sim. GBank gil3598974 (AF077000) - protein tyrosine phosphatase TD14 (Rattus norvegicus)	87651244 (5657, 5658) Novel Protein sim. GBank gij4680689jgbjAAD27734.1jAF13295 - (AF132959) CGI-25 protein [Homo sapiens]	88087109 (5659, 5660) Novel Protein sim. GBank gil2498667 sp Q61200 NPH1_MOUSE - NEUREXOPHILIN	2)	4)	87612938 (5665, 5666) Novel Protein sím. GBank gil5262615 emb CAB45747.1 - (AL080156) hypothetical protein [Homo sapiens]	18) Novel Protein slm. GBank gi[2224567 dbj BAA20772  - (AB002311) KIAA0313 [Homo sapiens]	87775712 (5669, 5670) Novel Protein sim. GBank gil 4589532 dbj BAA76788.1  - (AB023161) KIAA0944 protein [Homo sapiens]	85724748 (5671, 5672) Novel Protein sim. GBank gi[2351568 (U76618) - N-RAP
95320519 (5653, 5654)				87651244 (5657, 5658)		87614717 (5661, 5662)					_
2827			2828	2829	2830	2831	2832	2833	2834	2835	2836

	UNCLASSIFIED 18108394, 18108397, 264259, 29331826, 265007, 265019, 264448, 18108368, 20281149, 264565, 264567	UNCLASSIFIED 264555	264509, 264811, 265011, 264268, 264769, 265020, 264834, 264636, 264556	dna_rna_bind 56182575, 35696052, 264907, 264908, 264909, 264910, 264593, 264595, 264766, 265022, 264691, 33657182, 35695763, 18108370, 35695855, 264631, 264559, 264563, 264563	264600	UNCLASSIFIED 284908, 264907, 264908, 264910, 264764, 35695855, 83373044, 18108385	264685	homeobox 29146498, 87168474, 264686, 35696423, 83373044, 264564	UNCLASSIFIED 28331830, 264909, 265008, 265011, 8716859, 264629, 264556	UNCLASSIFIED 284908, 264766, 264769, 264629, 264637, 284566	UNCLASSIFIED 264490, 264259, 264508, 264905, 264907, 264510, 265007, 265008, 264591, 264592, 264592, 264595, 55812038, 265010, 265011, 264604, 264763, 264764, 264765, 264686, 264686, 264628, 264629, 264555, 264638, 264557, 264638, 264559, 264657, 264638, 264569, 264657, 264638, 264559, 264657, 264638, 264559, 264657, 264638, 264559, 264657, 264638, 264559, 264657, 264638, 264559, 264638, 264569, 264657, 264638, 264559, 264638, 264569, 264657, 264638, 264559, 264638, 264569, 264657, 264638, 264559, 264638, 264569, 264657, 264638, 264559, 264638, 264569, 264657, 264638, 264559, 264638, 264569, 264659, 264657, 264638, 264559, 264638, 264569, 264659, 264657, 264638, 264559, 264638, 264569, 264639, 264649, 2646499, 264649, 264649, 264649, 264649, 264649, 264649, 264649, 264649, 264649, 264649, 264649, 264649, 264649, 264649, 264649, 264649	29331822, 256960, 264509, 264906, 265007, 264594, 265018, 264288, 263972, 35696423, 18108384, 56526486, 18108390	UNCLASSIFIED 35696286, 265008, 265009, 265018, 264288,
				Contains protein domain (PF00076) - dna_rna_bind RNA recognition motif. (a.k.a. RRM, RBD, or RNP domain)			Contains protein domain (PF00008) - EGF-like domain				-		
2837   87766482 (5673, 5674) Novel Protein sim. GBank gil5420387 emb CAB46679.1  - (AJ243459) proteophosphoglycan [Lelshmania major]	87775392 (5675, 5676) Novel Protein sim. GBank gil973378 (U31263) - core protein [Hepatitis C virus]	85799317 (5677, 5678)	Novel Protein sim. GBank gij1575515 (U64899) - thrombospondin-related anonymous protein [Plasmodium gallinaceum]	86982568 (5681, 5682) Novel Protein sim. GBank gi 2224605 db  BAA20790  - (AB002330) KIAA0332 [Homo sapiens]		91012494 (5685, 5686) Novel Protein sim. GBank gil5578957 emb CAB51350.1  - (AL050306) d.1475B7.1 (novel KIAA0615 and KIAA0323  LIKE protein) [Homo sapiens]	56731154 (5687, 5688) Novel Protein sim. GBank gil585123 sp G08878 FBLC_MOUSE - FIBULIN-1, ISOFORM C PRECURSOR (BASEMENT-MEMBRANE PROTEIN 90) (BM-90)	94321719 (5689, 5690) Novel Protein sim. GBank gil5420387 emb CAB46679.1  -   (AJ243459) proteophosphoglycan (Lelshmania major)	88318613 (5691, 5692) Novel Protein sim. GBank (gi 5306263 gb AAD41995.1 AC00623 - (AC006233)   unknown protein [Arabidopsis thaliana]	81811757 (5693, 5694) Novel Protein sim. GBank gij3399676 (AC005390) - R31180_1 [Homo sapiens]	87612943 (5695, 5696) Novet Protein sim. GBank gil5262615 emblCAB45747.1  - (AL080156) hypothetical protein [Homo sapiens]	88084283 (5697, 5698) Novel Protein sim. GBank gij3342218 (AC004131) - Unknown gene product [Homo sapiens]	
87766482 (5673, 5674)			87774665 (5679, 5680)	86982568 (5681, 5682)	80080086 (5683, 5684)	91012494 (5685, 5686)							0 (87623636 (5699, 5700)
2831	2838	2839	284(	2841	2842	284:	3844	2845	2846	2847	2848	2849	<b>7820</b>

2851	87820548 (5701, 5702)	2851 87820548 (5701, 5702) Novel Protein sim. GBank gl 4321619 gb AAD15788.1  - (AF051098) seven transmembrane domain orphan receptor [Mus musculus]		UNCLASSIFIED	264906, 264907, 56182435, 264758, 55811386, 265010, 18108351, 264448, 264369, 21905765, 284691, 264692, 284693,
2852		86987023 (5703, 5704) Novel Protein sim. GBank gil1825729 (U88308) - similar to drosophila membrane protein PATCHED SP:P18502 (PID:g129645) [Caenorhabditis elegans]		UNCLASSIFIED	264591
	87784630 (5705, 5706)	87784630 (5705, 5706) Novel Protein sim. GBank gil2702347 (AF027503) - putative membrane-associated guanylate kinase 1 [Mus musculus]	Contains protein domain (PF00397) - kinase WW domain	kinase	56182575, 55811150, 264690, 27486262, 27485265, 264632, 56182323, 56526486, 2272000
2854	8808357 (5707, 5708)	88083557 (5707, 5708) Novel Protein sim. GBank gi[2795825 (AC004021) - ketch protein; ring canal component involved in cytoplasmic bridges; 77% Similarity to A45773 (PID:g1079096) [Homo sapiens]	Contains protein domain (PF01344) - dna_ma_bind Kelch motif	dna_ma_bind	35696286, 29331824, 29331826, 29331828, 264908, 264768, 264693, 22279002, 264482
	94723856 (5709, 5710)	94723856 (5709, 5710) Novel Protein sim. GBank gil1504040 dbj BAA13219  - (D86983) similar to D.melanogaster peroxidasin(U11052) [Homo sapiens]	Contains protein domain (PF00560) - glycoprotein Leucine Rich Repeat	glycoprotein	22278994, 29331822, 29331824, 29331825, 264906, 264908, 265009, 33109954, 265018, 265019, 265019, 2466265, 83373044, 22279000, 22279002, 26482
2856	88093359 (5711, 5712)	88093359 (5711, 5712) Novel Protein sim. GBank gij3264583 (AC005189) - match to ESTs H97758 (NID:g1118643) and AA085546 (NID:g1628773) [Homo sapiens]			21906766, 22278997, 265022, 29331822, 29331826, 27486262, 265007, 265009, 265017, 284482, 284563, 18108351
	95348286 (5713, 5714)	95348286 (5713, 5714) Novel Protein sim. GBank gi 3041855 (AC004537) - similar to tumor suppressor p33ING1; similar to AF044076 (PID:g2829208) [Homo sapiens]	Contains protein domain (PF00628) - struct PHD-finger	struct	22278995, 35696286, 29331824, 29331825, 35696052, 264103, 264108, 56182435, 21906765, 21906769, 265020, 18108368, 35695763, 22279002, 264563
2858	87434748 (5715, 5716)		Contains protein domain (PF00097) - dna_rna_bind Zinc finger, C3HC4 type (RING finger)	dna_rna_bind	264569, 264887, 22278995, 22278996, 22278996, 22278997, 22278997, 22278999, 264269, 29331826, 29331826, 29331827, 29331828, 264269, 264909, 264909, 264511, 264512, 265007, 265008, 264910, 265009, 264593, 60433356, 264596, 265018, 264764, 265009, 265010, 265011, 26469, 264764, 264765, 264768, 264769, 264769, 264769, 264769, 264769, 264769, 264697, 264692, 18108357, 264534, 264769, 264692, 18108370, 264692, 18108381, 264692, 18108381, 264692, 264685, 264566, 264568, 264568, 264686, 264568, 264686, 264568, 264568, 264686, 264568, 264686, 264566, 264566, 264566, 264566, 264566, 264486, 284567
6697	80937675 (5717, 5718)	8083/6/5 (5/11, 5/18) Novel Protein sim. GBank gild325320 gb AAD17331.1  - (AF124427) claudin-15 [Mus musculus]		UNCLASSIFIED	60424179, 65274572, 29331828, 264905, 264511, 264758, 265011, 21908767, 21906769, 55811957, 265021, 56182323

UNCLASSIFIED   CONTAINS   CONTA		<del>,</del>			,		, —	
gil4469186 emb CAB38414.1  - (novel protein similar to C. terial tRNA (5- indylate)-Methyltransferases)  gil3941730 (AF108083) - BS4 gil3941730 (AF108083) - BS4 gil3249088 (AC004473) - Contains protein domain (PF00096) - In protein gb M97204 from D. Zinc finger, C2H2 type thaliana] gil4078451 pir A55463 - cde - chicken gil43840 (L19048) - MSA-2 gil585703 sp Q07066 PMP2_RAT EMBRANE PROTEIN Lisculus]	264259, 29331822, 29331824, 29331825, 29331826, 29331827, 29331828, 264510, 264511, 33109954, 18108351, 264683, 264681, 264692, 264686, 21906765, 264691, 264692, 264693, 18108388, 22278002, 264882	264369, 264692	264691, 264638	56994075, 35696286, 22276998, 28331822, 29331824, 3569652, 29331828, 264106, 264511, 55812038, 33657084, 55811386, 255018, 265019, 21906765, 21906766, 21906769, 3365709, 255022, 33657023, 33657109, 3365739, 264629, 18108376, 60431650, 5618223, 18108376, 87168518, 22725007	264259, 264910	22278995, 21906764, 264482	65274572, 22278998, 22278998, 22278999, 264259, 29331824, 29331827, 29331828, 60433438, 21906754, 265018, 264448, 264764, 265018, 264448, 264764, 265018, 264448, 264764, 265018, 264448, 264764, 265018, 264448, 264764, 265018, 264448, 264764, 265018, 264448, 264764, 265018, 264448, 264764, 265018, 264448, 264764, 265018, 264448, 264764, 265018, 264448, 264764, 265018, 264764, 264764, 264764, 265018, 264448, 264764, 265018, 264448, 264764, 265018, 264764, 2647	264768, 264768, 21906768, 22278998, 265022, 264259, 264508, 264907, 265511, 264910, 264635, 264905, 264907, 265011, 265017, 285018, 265019, 264563, 264566, 264764, 264369, 264567, 264388, 264588, 264764, 264369, 264567,
gil4469186 emb CAB38414.1  - (novel protein similar to C. terial tRNA (5- indylate)-Methyltransferases) gil3941730 (AF108083) - BS4 gil3249088 (AC004473) - h protein gb M97204 from D. thaliana) thaliana) gil4658463 - chicken gil438840 (L19048) - MSA-2 gil585703 sp Q07056 PMP2_RAT	UNCLASSIFIED		UNCLASSIFIED	struct	struct		glycoprotein	UNCLASSIFIED
				Contains protein domain (PF00096) - Zinc finger, C2H2 type				
지 되는 그리 꼬리 이 있는 이번 있다. 그런 그리 그리	2860 87532599 (5719, 5720) Novel Protein sim. GBank gil4469186jembjCAB38414.1  - (AL031588) d.1163J1.2.1 (novel protein similar to C. elegans B0035.16 and bacterial IRNA (5- Melthylaminomethyl-2-thiouridylate)-Melthyltransferases) ((soform 1) [Homo sapiens]	18507 (5721, 5722) Novel Protein sim. GBank gij3941730 (AF108083) - BS4 [Homo sapiens]	99585 (5723, 5724) Novel Protein sim. GBank gil4505013 ref NP_002310.1 pLRN  - leucine-rich neuronal protein	:0421 (5725, 5726) Novel Protein sim. GBank gij3249088 (AC004473) - Contains sImilarity to goliath protein gb M97204 from D. metanogster. [Arabidopsis thaliana]	20030 (5727, 5728) Novel Protein sim. GBank gij1079451 pir  A55463 - tropomodulin, skeletal muscle - chicken	[2191 (5729, 5730) Novel Protein sim. GBank gi 438840 (L19048) - MSA-2 [Plasmodium falcipanum]	)5480 (5731, 5732) Novel Protein sim. GBank gij565703jspjQ07056jPMP2_RAT - 22 KD PEROXISOMAL MEMBRANE PROTEIN	98001 (5733, 5734) Novel Protein sim. GBank gil4580997lgbJAAD24571.1JAF12108 - (AF121081) cAMP inducible 2 protein [Mus musculus]

18108392, 264488, 22278994, 22278997, 22278998, 60432049, 264259, 29331822, 29331824, 29331825, 60432289, 29331822, 29331827, 35686052, 29331828, 264508, 52644045, 264828, 265008, 265009, 264591, 60432229, 264507, 265009, 264591, 60432229, 264593, 60433356, 264595, 21906754, 265017, 265019, 264595, 21906754, 21906765, 264563, 264563, 264564, 264567, 264487	264369	264905, 264908, 264764, 21906769, 264634	264259, 29331822, 60432289, 29331827, 264907, 265008, 265017, 265018, 264682, 264764, 18108354, 265021, 27486265, 264629, 18108387, 264567	263981	264488, 18108374, 264768, 264687, 264688, 264689, 35696423, 35696286, 3569517, 264510, 264511, 265007, 264512, 265008, 264510, 264514, 265007, 264512, 265008, 264590, 264534, 264634, 264555, 264592, 264559, 264536, 60433289, 35696052, 265011, 264600, 264509, 264509, 264508, 264508, 264563, 18108370, 264507, 264566, 264908, 264764, 264288, 264567, 264909, 264766, 18108391
·	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	kinase	UNCLASSIFIED
2868   95303283 (5735, 5736)   Novel Prolein sim. GBank gil 1292868 emb CAA63923  (X94232) 1-Cell activation protein [Homo sapiens]			2871   88318621 (5741, 5742) Novel Protein sim. GBank gi 5306263 gb AAD41995.1 AC00623 - (AC006233) unknown protein [Arabidopsis thaliana]	95312197 (5743, 5744) Novel Protein sim. GBank gil112205 pir  B39066 - proline- inch protein 15 - rat	
95303283 (5735, 5736)	88094412 (5737, 5738)	84404574 (5739, 5740)	88318621 (5741, 5742)	95312197 (5743, 5744)	88094252 (5745, 5746) ,
2868	2869	2870	2871	2872	2873

52644507, 52645156, 52646365, 56182575, 22278994, 22278994, 22278995, 56994075, 35696286, 60432049, 264259, 52695080, 29331822, 56182181, 29331824, 60424269, 29331825, 66714117, 29331824, 60424269, 29331825, 26331828, 33656970, 60712502, 52644045, 268007, 265009, 60433356, 284758, 2357084, 265017, 264604, 265018, 265619, 264682, 264369, 264604, 265019, 264604, 265012, 265020, 265021, 265022, 52644150, 33657023, 52645129, 18108374, 18108376, 33657023, 52645129, 18108387, 87168518, 60432113, 22279000, 22279002, 264563, 264565	22278996, 22278997, 22278999, 29331826, 29331828, 29331828, 29146499, 66712502, 265008, 265017, 18108351, 264683, 264689, 21906767, 18108376, 18108377, 55811576, 60170394, 22279000, 264487		52646365, 65274572, 56182575, 22278997. 22278998, 22278999, 60432049, 52645080. 60424269, 60432289, 29331827, 35696052, 29331828, 66712502, 52644045, 56182435, 60433358, 33657402, 33657084, 265019, 55811150, 264448, 264369, 21906766, 21906768, 21906769, 265020, 33657023, 33657109, 33657182, 27486262, 264629, 60431528, 55811576, 52644332, 56182323, 264558, 83373044, 18108385, 56526486, 6045113, 22279000, 22279002, 264563	264905, 264509, 2533 1022, 2533 1020, 264905, 264509, 264907, 264909, 264510, 265007, 264907, 264909, 264750, 21905764, 18108351, 264681, 264681, 264688, 264688, 264688, 264689, 264689, 264693, 264693, 18108364, 33657109, 264629, 18108374, 264630, 264632, 264653, 264653, 264563, 264563, 264563, 264563, 264563, 264563, 264563, 264565, 264566, 264486, 264567
UNCLASSIFIED	glycoprotein	igi		ONCLASSITED
	Contains protein domain (PF00560) - glycoprotein Leucine Rich Repeat	Contains protein domain (PF00167) - ígf Fibroblast growth factor		Contains protein domain (Pr01207) - UNCLASSIFIED Uncharacterized protein family UPF0034
2874   94313549 (5747, 5748) Novel Protein sim. GBank gij3212854 (AC004005) - unknown protein [Arabidopsis thaliana]	88083726 (5749, 5750) Novel Protein sim. GBank gil2781386 (AC004010) - similar to Leucine-rich transmembrane proteins: 44% similarity to U42767 (PID:g1736918) [Homo sapiens]	88090854 (5751, 5752) Novel Protein sim. GBank gil2979530 (AC004449) - R33683 2 [Homo sapiens]	94747029 (5753, 5754) Novel Protein sim. GBank gil4704208 emb CAB41646.1  - (AL035419) dJ1100H13.1 (putative novel protein) [Homo sapiens]	88095309 (5755, 5756) Novel Protein sim. GBank gij3876775jembjCAB03067j- (Z81077) predicted using Genefinder, Similarity to Yeast protein 8248 (TR:G587531) [Caenorhabditis elegans]
4 94313549 (5747, 5748)	2875 88083726 (5749, 5750)	2876 88090854 (5751, 5752)		2878   88095309 (5755, 5756)

				lentrecept 60432049, 264259, 29331828, 264908. 264511, 264595, 60433438, 264598, 265017, 264605, 283969, 283972, 264555, 83373044, 87168518, 264566 SIFIED 18108351, 264686, 264629, 264631, 264639, 83373044, 284482 R3373044, 284482	
Contains protein domain (PF00315) - UNCLASSIFIED Uracil-DNA glycosylase		UNCLASSIFIED	Contains protein domain (PF00400) - UNCLASSIFIED WD domain, G-beta repeat UNCLASSIFIED	Contains protein domain (PF00084) - complementrecept Sushi domain (SCR repeat) UNCLASSIFIED Contains protein domain (PF00098) - transcriptfactor	Size fine fineer. C2H2 type
2879   87869122 (5757, 5758)   Novel Protein sim. GBank gi 4895145 gb AAD32752.1  - (AF127374) unknown [Streptomyces lavendulae]	94851439 (5759, 5760) Novel Protein sim. GBank gij4880703jgbjAAD27741.1jAF13296 - (AF132966) CGI-32 protein [Homo sapiens]	87650539 (5761, 5762) Novel Protein sim. GBank gij733571 (U23452) - No definition line found [Caenorhabdilis elegans] 87714367 (5763, 5764) Novel Protein sim. GBank gij1118112 (U41559) - No definition line found [Caenorhabdilis elegans]	95362875 (5765, 5766) Novel Protein sim. GBank 914868008lgb/AAD31087.1JAF10593 - (AF105934) vitamin D receptor-interacting protein [Homo sapiens] 87784643 (5767, 5768) 83006308 (5769, 5770) Novel Protein sim. GBank gij2224697[dbjjgAA20832] - (AB002376) KIAA0378 [Homo sapiens]	91237823 (5771, 5772) Novel Protein sim. GBank gi[1255889 (U53344) - T07H6.5 gene product [Caenomabditis elegans] 91227860 (5773, 5774) Novel Protein sim. GBank gi[3882323 db] BAA34521.1  - (AB018344) KIAA0801 protein [Homo sapiens] 95105816 (5775, 5776) Novel Protein sim. GBank	dil4508015 refine 2015   Contain State   Contains protein 2015   Contains protein 2015   Contains protein 2015   Contains State   Contains Sta
78 87869122 (5757, 5758)		2881 87650539 (5761, 5762) 2882 87714367 (5763, 5764)	2883 95362875 (5765, 5766) 2884 87784843 (5767, 5768) 2885 83006308 (5769, 5770)	2886 91237823 (5771, 5772) 2887 91227860 (5773, 5774) 2888 95105816 (5775, 5776)	

2889	87606562 (5777, 5778)			UNCLASSIFIED	56994075, 29331824, 265009, 264760,
2880	78703853 (5779, 5780)	78703853 (5778, 5780) Novel Protein sim. GBank gil854065 emb CAA58337  - (X83413) U88 [Human herpesvirus 6]		UNCLASSIFIED	264591, 264766
2891	88094428 (5781, 5782)	88094428 (5781, 5782) Novel Protein sim. GBank gij3877750jembjCAB01508j - (Z78084) predicted using Genefinder; similar to collagen; cDNA EST EMBL:D65885 comes from this gene; cDNA EST EMBL:D69451 comes from this gene; cDNA EST EMBL:D6026 comes from this gene; cDNA EST EMBL:D69658 comes from this gene; cDNA EST		UNCLASSIFIED	264628, 264593, 284369, 264685, 264693, 264628, 264563, 284566
2892	95419745 (5783, 5784)	95419745 (5783, 5784) Novel Protein sim. GBank gil4929759jgb AAD34140.1 AF15190 - (AF151903) CGI-145 protein [Homo sapiens]		UNCLASSIFIED	65274572, 35696286, 29331828, 264110, 265009, 60433438, 265018, 265019, 18108351, 264288, 21906765, 21906766, 21906789, 55811957, 264690, 65274620, 263967, 35695855
2893	87798014 (5785, 5786)			UNCLASSIFIED	264259, 265010, 264682, 18108370, 264555, 264556
2894		87755985 (5787, 5788) Novel Protein sim. GBank gij5669015 gb AAD46135.1 - (AF080171) zinc finger protein ZNF232 [Homo sapiens]	Contains protein domain (PF00096) - Irranscriptfactor Zinc finger, C2H2 type	transcriptfactor	264259, 265006, 60433438, 52644296, 265011, 264369, 35695917, 18108381, 18108382, 18108388
2895		86938778 (5789, 5790) Novel Protein sim. GBank gil3924708 emb CAA84646  - (Z35597) Weak similarity with sea squirt nidogen precursor protein (blastp score 71); cDNA EST EMBL:T02069 comes from this gene; cDNA EST EMBL:D76135 comes from this gene; cDNA EST EMBL:D73147 comes from this gene; cDNA EST EMBL.D73147 comes from this gene;	Contains protein domain (PF01437) - Plexin repeat		29331824, 265007, 264762, 264636, 264563
2896		87752122 (5791, 5792) Novel Protein sim. GBank gl 4885549 ref NP_005456.1 pPKBG - protein kinase B gamma	Contains protein domain (PF00069) - kinase Eukaryotic protein kinase domain	kinase	78108392, 18108394, 18108398, 22278998, 264259, 29331824, 29146499, 264906, 265007, 265009, 265018, 265019, 264685, 264689, 21906766, 265021, 264693, 33657182, 264839, 18108384, 18108388, 264567
2897		95413057 (5793, 5794) Novel Protein sim. GBank gil4502977 refiNP_001296.1 pCLDN - Clostridium perfringens enterotoxin receptor 1		UNCLASSIFIED	60424179, 56182575, 22278996, 35696286, 22278997, 22278999, 60432049, 264259, 29331824, 29331825, 60424269, 60432289, 29331826, 29331826, 29331828, 35696052, 264908, 56182435, 265009, 264910, 60170831, 60431735, 60433356, 60433438, 65274444, 55811386, 265018, 18108351, 264448, 264288, 264687, 21906765, 21906766, 21906762, 21906764, 264534, 33657023, 33657109, 35695763, 264528, 264628, 264629, 60431528, 18108374, 25810764, 55811576, 35696423, 35696855, 264555, 56182323, 18108385, 264404, 22279000, 22279000, 22279000, 2264586
2898	87750340 (5795, 5796)			UNCLASSIFIED	22278995, 264604, 18108385, 264566

2899				Γ	000000000000000000000000000000000000000
<u>8</u>	8035/6/0 (5/8/, 5/88)		וכ	3	204704, 21900704, 264092
	94233538 (5789, 5800)	2900  94233538 (5799, 5800) Novel Protein sim. GBank gij4581470 emb CAB40137.1 -	<u> </u>	glycoprotein	65274572, 56182575, 35696286, 60432049,
		(Y18483) SLC7A8 protein [Homo sapiens]			264259, 29331824, 66714117, 29331826,
_					35696052, 29331828, 66712502, 56182435,
_					265006, 265007, 265008, 265009, 60433356,
					264758, 265018, 264764, 264765, 264288,
					264768, 21906764, 21906768, 21906769,
					265020, 264692, 264693, 32833986, 264631,
					83373044, 60432113
2901 8	87444731 (5801, 5802)	87444731 (5801, 5802) Novel Protein sim. GBank	<u>a</u>	phosphatase	22278995, 22278997, 22278999, 60432049,
		gil4759272 ref NP_004614.1 pTTC4 - tetratricopeptide			29331822, 29331824, 29331825, 29331827.
_		repeat domain 4			35696052, 33656970, 284910, 265009,
_					21906754, 33657084, 87168474, 265010.
_					265018, 21906764, 21906765, 21906766,
					21906767, 21906769, 33657023, 264693,
					33657109, 33657349, 35696423, 35695855,
-					263981, 56182323, 22279002
2902	85745271 (5803, 5804)	85745271 (5803, 5804) Novel Protein sim. GBank gij2414615jembjCAB16364j -			264683, 264691
		(299259) hypothetical protein (Schizosaccharomyces			
		ротреј			
2903	87606733 (5805, 5806)	87606733 (5805, 5806) Novel Protein sim. GBank gi 1079318 pir  S52241 - XLCL2			264887, 22278994, 264259, 29331826,
		protein - African clawed frog			29331828, 264905, 52644045, 56182435,
					264511, 265017, 265018, 18108351, 264448,
					264683, 264769, 264689, 35695917,
_					52644150, 87168518, 60432113, 22279002
2904	86458072 (5807, 5808)	86458072 (5807, 5808) Novel Protein sim. GBank	7	UNCLASSIFIED	52646365, 22278999, 264259, 35696052,
		gi[5639823[gb[AAD45885.1[AF14367 - (AF143676)			265011, 265017, 264683, 264769, 35695917,
		multispanning nuclear envelope membrane protein nurim			265020, 263967, 18108374, 35695855.
		[Homo sapiens]			264637, 264952, 18108385, 18108387
2905	84449926 (5809, 5810)	84449926 (5809, 5810) Novel Protein sim. GBank		oncogene	265009, 264681, 264682
_		gij728837 spjP39194 ALU7_HUMAN - !!!! ALU SUBFAMILY			
		SQ WARNING ENTRY IIII			
3008 2808	95341051 (5811, 5812)			UNCLASSIFIED	22278996, 35696286, 22278998, 264259,
		gi[4689256]gb[AAD27831.1[AF12185 - (AF121858) sorting			60432289, 29331828, 29331830, 66712502,
_		nexin 8 [Homo sapiens]			265009, 60170831, 33109954, 264448,
_					264683, 264288, 264689, 21906766,
					21906767, 21906768, 55811957, 35695917,
					265022, 52644150, 264691, 33657023,
					264692, 264693, 35695855, 60432113,
1					264566

2907	91211383 (5813, 5814)	gil1707079 (U80451) - contains	Contains protein domain (PF00226) - eph	ųd:	52644507, 56182575, 56181686, 22278995, 5694075, 35696286, 60432049, 56182181,
	_	Strong similarity to a Cives-like dolinality (19.1 Society) [Caenorhabditis elegans]			35696052, 60431735, 264595, 55812038.
					21906754, 55811386, 265019, 264682,
					264369, 56181562, 21906766, 55811957,
		•			35695917, 265020, 265021, 33657023,
					33657109, 60431528, 55811576, 35696423.
					35695855, 264638, 22279000
2008	_	80414246 (5815 5816) Novel Protein 8im (5Bank pii2673917 (AC002561) - putalive		helicase	265009, 33109954, 18108351, 264766.
200	לפוסה יהוסה) סרידרו הסי	ATP-dependent RNA helicase [Arabidopsis thaliana]			265021, 264691, 264692, 18108374, 264556,
					264638, 264557, 264558
2000	87420225 (5817 5818)			eph	264259, 87168474, 265018, 18108365.
3					264628
2910	86601075 (5819 5820)	86601075 (5819 5820) Novel Protein sim. GBank gil4539335lemblCAB37483.11-			22278995, 264509, 264512, 265007,
	, , , , , , , , , , , , , , , , , , , ,				33657402, 265017, 264369, 265022,
		formers and some district and formers			18108365, 264628
2011	7	94216615 (5821 5822) Navel Protein sim GBank pil4469187lembICAB38415.11-	5	glucoamylase	52646365, 18108397, 22278995, 22278997,
	(				22278998, 22278999, 29331824, 29331825.
		Root Home saniens			52844045, 265008, 265018, 264448,
					21906765, 21906766, 21906767, 21906768,
	-				21906769, 55811957, 265021, 18108370,
					18108372, 18108374, 22279000
2912	87731803 (5823 5824)	87731803 (5823, 5824) Novel Protein sim, GBank	Contains protein domain (PF00904) -		52645156, 264092, 60432049, 264259,
!		gil4929637[qb[AAD34079,1]AF15184 - (AF151842) CGI-84 Involucin repeat	Invotucrin repeat		52645080, 29331824, 29331825, 66712502.
		protein (Homo sapiens)			33109954, 264760, 264683, 264288, 264686,
					265021, 264693, 18108368, 263976, 264404
2913	87713823 (5825, 5826)	87713823 (5825, 5826) Novel Protein sim. GBank gil854065lemb CAA58337  -	Contains protein domain (PF00096) - UNCLASSIFIED	UNCLASSIFIED	52644507, 52645156, 52646842, 56182575,
		(X83413) U88 [Human herpesvirus 6]	Zinc finger, C2H2 type		35696286, 22278997, 264259, 52645080,
	•				29331827, 35696052, 29331828, 264828,
					52644045, 56182435, 55812038, 52646317.
					21906754, 52644296, 87168474, 265017.
					265018, 265019, 18108351, 264682, 264686,
					264689, 21906765, 21906766, 21906767.
					21906768, 35695917, 265020, 52644150,
					27486261, 27486262, 27486265, 35695763.
	-				55811576, 35695855, 52644332, 22279000.
					22279002, 264563
2914	2914 87797300 (5827, 5828)				264557

	98081972 (5829, 5830)	98081972 (5829, 5830) Novel Protein sim. GBank gi 5174485 ret NP_006030.1 pKIAA - endocytic receptor (macrophage mannose receptor family)	Contains protein domain (PF00059) - eph Lectin C-type domain	· ·	264569, 264488, 264687, 264768, 21906766. 52646842, 21906767, 21908768, 56182575, 29148629, 35695917, 22278996, 22278997, 22278998, 22278997, 22278998, 25278997, 22278998, 25644150. 264691, 264259, 60432049, 264692, 25486261, 35696052, 29331828, 27486262, 27486262, 29331828, 27486262, 293657049, 29146498, 264906, 264907, 18108374, 265008, 60432229, 264593, 60433356, 265008, 60432229, 264593, 60433356, 56182323, 18108382, 33657084, 87168518, 265010, 265011, 60432113, 265017, 265018, 22279000, 265019, 55811150, 264681, 18108354, 264388, 264488, 264683, 264566, 18108354, 264389, 264488, 264683, 264566, 18108354, 264389, 264488, 264683, 264566, 18108354, 264389, 264488, 264683, 264566, 18108354, 264389, 264488, 264683, 264566, 18108354, 264389, 264488, 264683, 264566, 18108354, 264389, 264488, 264683, 264566, 18108354, 264389, 264488, 264683, 264566, 18108354, 264388, 264588, 264588, 264388, 264566, 18108354, 264388, 264488, 264683, 264566, 18108354, 264388, 264488, 264683, 264566, 18108354, 264388, 264488, 264683, 264566, 18108354, 264388, 264488, 264583, 264566, 18108354, 264388, 264488, 264583, 264566, 18108354, 264388, 264488, 264583, 264566, 18108354, 264388, 264488, 264583, 264566, 18108354, 264388, 264488, 264583, 264566, 18108354, 264488, 264488, 264583, 264566, 18108354, 264388, 264488, 264583, 264566, 18108354, 264388, 264488, 264583, 264566, 18108354, 264388, 264488, 264583, 264583, 264583, 264584, 264584, 264583, 264584, 264583, 264584, 264583, 264584, 264584, 264584, 264583, 264584, 2
		95337790 (5831, 5832) Novel Protein sim. GBank gij5104851 dbjjBAA80165.1 - (AP000061) 305aa long hypothetical dTDP-4- dehydrorhamnose reductase [Aeropyrum pernix]		dehydrogenase	5245156, 65274572, 22278994, 22278995, 35696288, 22278996, 22278997, 22278998, 22278998, 22278999, 22278998, 22278999, 22278998, 22331824, 29331825, 66714117, 60432289, 29331826, 29331827, 29331820, 29331827, 29331830, 5264056, 264509, 26182435, 60170831, 264592, 264509, 265011, 265017, 265018, 26644296, 85658542, 265011, 265017, 265018, 265019, 18108351, 26448, 264288, 5264429, 21906765, 21906765, 21906766, 21906766, 21906766, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906765, 21906769, 265022, 27466261, 27466262, 27466264, 35695763, 18108376, 55811576, 35696423, 6527791, 26508323, 18108376, 35696423, 62279002, 264482
		87454546 (5833, 5834) Novel Protein sim. GBank gij3169065 emb CAA19260.1  - (AL023704) putative translocation elongation factor-Tu family (Schizosaccharomyces pombe)	. 4	UNCLASSIFIED	60433438, 264602, 264682, 87168518. 60432113
		85690528 (5835, 5836) Novel Protein sim. GBank gij539218 pir  S38038 - hypothetical protein YKL201c - yeast (Saccharomyces cerevisiae)			264638
2919	87641497 (5837, 5838)	87641497 (5837, 5838) Novel Protein sim. GBank gi 2564955 (AF030001) - unknown [Mus musculus]			66714117, 66712502, 263981

(0+0) (000)			250331824, 29331826, 29331827, 264828,
			264909, 26182435, 264211, 264738. 33109954, 21908754, 52644296, 265010.
			285011, 264601, 265017, 265019, 264681,
			264690, 264691, 264692, 264693, 33657109.
			33657182, 27486262, 27486264, 27486265,
			35696423, 35695855, 264632, 264636,   264637   264638   66182323   60170394
			18108385, 87168518, 60432113
91639982 (5841, 5842) Novel Protein sim. GBank	Contains protein domain (PF00787) -		35696286, 22278997, 264091, 264092,
	PX domain		264094, 264259, 29331822, 29331824.
associated factor 2 [Homo sapiens]			29331826, 29331827, 35696052, 29146498, 264104 264105 264107 264509 264110
			264112, 264512, 60433356, 21906754,
		•	87168474, 265017, 18108351, 264288.
			21906765, 21906766, 21906767, 21906769,
			35695917, 265021, 263974, 18108374,
			263976, 263977, 18108376, 264555, 263981,
			56526486, 87168518, 22279000, 22279002
87749762 (5843, 5844) Novel Protein sim. GBank gil4589514 ldbjjBAA76779.1  -	Contains protein domain (PF01074) - kinase	kinase	264906, 264909, 264511, 265006, 265008,
(AB023152) KIAA0935 protein [Homo sapiens]	Glycosyl hydrolases family 38		264593, 33657402, 60174639, 18108351,
ι			264/63, 219U6/65, 2914862/, 3509591/,
			264692, 264629, 263978, 55811576,
			60170394 22279000 264486
95337799 (5845, 5846) Novel Protein sim. GBank gil4835268lembiCAB42898 21 -	Contains protein domain (PF00169) - struct	struct	264488, 18108397, 22278995, 22278996.
	PH domain		22278997, 22278998, 22278999, 29331825,
[Homo sapiens]			29331826, 29331827, 29331830, 264511.
			265009, 33657402, 265011, 265017, 265018,
			264683, 18108354, 21906765, 21906767,
			21906768, 21906769, 52644150, 264691,
			264692, 33657109, 263974, 18108376,
			264631, 264636, 18108385, 18108387,
			22279000, 264563, 264568
87791957 (5847, 5848)   Novel Protein sim. GBank gi 2133095 pir  572254 -  riberomal protein   36 milespondial   1967	Contains protein domain (PF00444) - ribosomalprot	ribosomalprot	265017, 264628, 20281152, 264556
(Saccharomyces cerevisiae)	Aibusoinal protein Loo		
95090120 (5849, 5850) Novel Protein sim. GBank gil2388986jemb CAB11718] -		UNCLASSIFIED	56182575, 35686286, 264259, 60432289,
(298980) actin associated protein [Schizosaccharomyces			29331827, 264508, 52644045, 264910.
[equod			264591, 60432229, 55812038, 21906754,
			264681, 264448, 264683, 264288, 264585,
	•		52644229, 264689, 21906/65, 21906/66,
			60170614 264602 3365702 26362.
			10010, 10001, 10001, 100000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 100000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 100000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 100000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 100000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 10000, 1000

	95343003 (5851, 5852)				29331828 265011 264768 264689
2927	80408018 (5853, 5854)	80408018 (5853, 5854) Novel Protein sim. GBank gi[283032 pirj 522456 - hvdrovyntoline-rich olymonoline i parancial taneluta			264764, 264288, 264630, 264637
	20452179 (5855, 5856)	20452179 (5855, 5856)		INCI ASSIFIED	264559
2929	91622920 (5857, 5858)	Novel Protein sim. GBank gil34133201emblCAA069151 -		INCI ASSIFIED	254550 254480 22278004 35505285
		(AJ006215) CMP-N-acetylneuraminic acid synthetase fMus			22278006 22278008 22278000 264004
		musculus			264259 52645080 29331822 29331824
			r	•	66714117, 29331825, 29331826, 29331827,
					35696052, 33656970, 264109, 29331830,
					52844045, 265009, 33109954, 52644296,
					87168559, 264760, 264762, 264448, 264764,
					264288, 264766, 264768, 21908765,
					21906766, 21906768, 21906769, 35695917.
					264691, 33657023, 264693, 33657109,
					18108374, 263976, 35696423, 35695855,
					263981, 22279000, 22279002, 264567,
					264486
2830	95302755 (5858, 5860)			UNCLASSIFIED	56182575, 56181686, 35696286, 22278996,
					22278998, 22278999, 264259, 29331825,
					60432289, 29331828, 264905, 52644045,
					56182435, 265009, 60170831, 264592,
					60432229, 60433356, 87168474, 265010,
					265011, 265017, 265018, 265019, 264762,
					264448, 264683, 264288, 264766, 21906765,
					21906769, 35695917, 60170615, 33657023,
					33657109, 264628, 18108370, 18108372,
					35696423, 35695855, 264556, 56182323,
, 600					60432113, 264567
1887	94312693 (5851, 5862)	94312693 (3851, 5862) Novel Protein sim. GBank gij3786433 (AF098505) - similar	Contains protein domain (PF00471) - UNCLASSIFIED	UNCLASSIFIED	52645156, 22278997, 22278998, 29331822,
		to Arabidopsis thallana male sterllity protein 2 (SW:Q08891) Ribosomal protein L33	Ribosomal protein L33		52645080, 29331824, 60432289, 33656970.
		[Caenorhabditis elegans]			60433356, 60433438, 33109954, 21906765.
					21906766, 21906767, 21906768, 265020,
					52644150, 33657023, 33657109, 33657182,
_					27486265, 35698423, 35695855, 264555,
20.00	7003 6000 6000				87168518, 60432113, 264566
7 6	19022023 (3603, 3604)				264906, 264907
5567	9172070 (3865, 3866)	91720770 (3865, 3865) Novel Protein sim. GBank gij3378056 (AF017777) - helicase		helicase	264488, 18108392, 56182575, 22278999,
		Losophila melanogasteri			264091, 264259, 29331825, 60432289.
					29331827, 264508, 52644045, 56182435,
			-		265007, 265009, 264592, 60433356.
					60433438, 21906754, 265017, 264682,
					264288, 52644229, 21906765, 21906768.
					21908768, 21906769, 265022, 52644150.
					33657023, 33657109, 27486265, 264635,
		-			264636, 60170394, 56182323, 18108385,
					60432113, 264565, 264566, 264567

2934	86576025 (5867, 5868)				22278997, 22278999, 29331824, 33657402, 264691, 27486282, 264628, 87168518, 22279000
2935	86410579 (5869, 5870)			UNCLASSIFIED	56182575, 22278995, 60433356, 33657402, 264758, 33109954, 21906754, 265018, 265019, 264448, 264769, 21906764, 21906765, 265021, 264692, 33657023,
2936		87605863 (5871, 5872) Novel Protein sim. GBank gil4153862 (AC005065) - determined by GENSCAN prediction and spliced EST; match to EST R84329 (NID:942735) Homo saniens!	Contains protein domain (PF00856) - nuclease SET domain	nuclease	33697109, 33697,349, 35810764, 22278000 22278997, 29331827, 29331828, 265009, 265077, 264605, 265020, 55811576, 18108387, 60432113, 264663
2937	94853096 (5873, 5674)	94853096 (5873, 5874) Novel Protein sim. GBank gij5174409 ref NP_006101.1 pCD28 - CD2 antigen (cytoplasmic tail)-binding protein 2		UNCLASSIFIED	56994075, 22278999, 264259, 60432049, 29331822, 56182181, 29331827, 29331828, 264906, 264909, 56182435, 265006, 264512, 264910, 60170831, 60433358, 265011, 265018, 18108351, 26448, 264288, 264766, 52644229, 21906765, 29148784, 65274791, 26456, 66182232, 60170394, 26458, 26458, 26458, 26458, 26458, 26458, 264584, 2645
2938		95419773 (5875, 5876) Novel Protein sim. GBank gij3319990 emb CAA76720  - (Y17267) ubiquitin-conjugating enzyme [Mus musculus]	Contains protein domain (PF00179) - ubiquitin Ubquitin-conjugating enzyme		264488, 56182575, 22278996, 35696286, 22278997, 22278999, 264490, 264259, 29331824, 66714117, 29331827, 35696052, 264107, 264906, 262008, 25264045, 56182435, 264511, 265008, 255009, 60432229, 33657402, 66433438, 55812038, 21806754, 85658542, 265010, 285011, 87168559, 265017, 265018, 265019, 264681, 264680, 264691, 265019, 264681, 264690, 264691, 265020, 60170815, 264690, 264691, 264691, 265020, 60170815, 264690, 33657109, 18108370, 18108381, 26395855, 264556, 18108381, 5650848, 83373044, 18108385, 5650848, 83373044, 18108385, 5650848, 83373044, 18108385, 5650848, 83373044, 18108385, 56508488, 83373044, 18108385, 85508488, 83373044, 18108385, 85508488, 83373044, 18108385, 85508488, 83373044, 18108385, 85508488, 83373044, 18108385, 85508488, 83373044, 18108385, 85508488, 83373044, 18108385, 85508488, 83373044, 18108385, 85508488, 83373044, 18108385, 85508488, 83373044, 18108385, 85508488, 83373044, 18108385, 85508488, 83373044, 833740404, 83374040404, 83374040404, 83374040404040404040404040404040404040404
2939	87786622 (5877, 5978)	87786622 (5877, 5878) Novel Protein sim. GBank gij3979900[emb]CAA99909] - (Z75547) similar to WD domain, G-beta repeat; cDNA EST yk371b7.5 comes from this gene; cDNA EST yk46545.5 comes from this gene; cDNA EST yk46545.5 comes from this gene; cDNA EST yk472c4.5 comes from this gene; cDNA EST yk472c4.5 comes from this gene; cDNA EST yk472c4.5 comes from this gene;	Contains protein domain (PF00400) - ATPase_associated WD domain, G-beta repeat	ATPase_associated	

22278996, 29331822, 29331824, 66714117, 29331826, 29331828, 264905, 264908,	66712502, 29331830, 265011, 265017, 264764, 264359, 21906766, 21906767, 33657023, 33657109, 32833986, 18108374,	18108377, 264634, 83373044, 18108385, 18108387, 264566	57	264910, 265010, 264768	60432049, 264259, 60432289, 29331827, 29146498, 265008, 284593, 60433356, 60433438, 265010, 265011, 265017, 265018, 264683, 264766, 18108381, 65274727, 60432113, 264567	56994075, 22278938, 50432049, 224229, 29331822, 29331824, 60424269, 60432289, 29331824, 26424269, 60432289, 29331824, 26424269, 60432289, 25331824, 60424269, 60432289, 2564909, 264511, 265006, 265009, 264603, 264603, 265017, 265018, 18108351, 264682, 264660, 21906767, 21906768, 21906769, 265021, 60170615, 52644150, 264690, 264691, 33657023, 264692, 265693, 33657109, 33657102, 264635, 264556, 56182323, 60170394, 264558, 264559, 83373044, 264582, 264567, 264482, 264563, 264484, 264567	60424179, 22278995, 22278996, 22278998, 22278999, 22278999, 264259, 56182181, 29331824, 60424269, 2643229, 56182181, 29331824, 265006, 60433286, 55812038, 264759, 55811386, 265018, 264681, 18108351, 264813, 264687, 21906768, 21906769, 35695917, 265020, 265021, 264693, 60431528, 55810764, 35896423, 35695855, 264630, 80170394, 83373044, 22279000, 285656, 264567
UNCLASSIFIED 2227	6671 2647 3365		UNCLASSIFIED 284557	glycoprotein 2649	collagen 604; 2914 604; 2646 604:		
						Contains protein domain (PF00415) - ATPase_associated Regulator of chromosome condensation (RCC1)	Contains protein domain (PF00096) - UNCLASSIFIED Zinc finger, C2H2 type
			21423370 (5881, 5882) Novel Protein sim. GBank gij3413872jdbjjBAA32300  -  AB007924) KIAA0455 protein [Homo saptens]	87430203 (5883, 5884) Novel Protein sim. GBank gil 172845 sp P46629 RB25_RABIT - RAS-RELATED PROTEIN RAB-25	95314504 (5885, 5886) Novel Trotein sim. GBank gil4929653jgb AAD34087.1 AF15185 - (AF151850) CGI-92 protein [Homo saplens]	2944 95081063 (5887, 5888) Novel Protein sim. GBank gil4678282jembjCAB41190.1j - (AL049660) 1-acytcerol-3-phosphate acyttransferase-like protein [Arabidopsis thaliana]	94233560 (5889, 5890) Novel Protein sim. GBank gij728831jspjP39188jALU1_HUMAN - IIII ALU SUBFAMILY Zinc finger, C2H2 type J WARNING ENTRY IIII
2940  95011103 (5879, 5880)	-50-		21423370 (5881, 5882) N	87430203 (5883, 5884) N 91 91	95314504 (5885, 5886)	95081063 (5887, 5888) P	94233560 (5889, 5890)
2940			2841	2942	2843	29444	2945

	UNCLASSIFIED 22278995, 22278997, 22278997, 22278999, 22278999, 22278999, 22278997, 22278999, 22278999, 22278999, 229146498, 265008, 265008, 2630189, 265019, 265019, 265019, 265019, 265019, 264086, 21906768, 21906768, 21906768, 21906769, 264628, 18108370, 264629, 264629, 18108387, 60432113	52646842, 22278995, 264259, 29331824, 29331825, 29331827, 29331830, 264909, 265007, 265009, 265019, 264763, 264684, 264288, 264688, 264688, 264688, 264689, 18108374, 55811576, 18108385, 22279002, 264553, 264567	UNCLASSIFIED 22278998, 22278997, 60432289, 29331826, 29331827, 29331828, 35695052, 28146499, 264104, 264107, 264905, 66712502, 264908, 60433356, 60433438, 87168559, 264764, 52644229, 56181562, 21906767, 21906768, 21906769, 265022, 60170615, 33657023, 35696423, 263981, 284558, 60432113, 22279002		264693
5	5	*		Contains protein domain (PF00883) - peptidase Cytosol aminopeptidase family	
2946   94317315 (5891, 5892) Novel Protein sim. GBank gi 541952 gb AD43195.1 AF07286 - (AF072864) peroxisomal membrane protein PMP 24 [Homo sapiens]	87362952 (5893, 5894) Novei Protein sim. GBank gil3540281 gb AAC34383.1  - (AF056116) All-1 related protein [Fugu rubripes]	87626527 (5895, 5899) Novel Protein sim. GBank gil5566014 gb AAB65654.2  - (AF001533) mitogen-induced [Mus musculus]	88175545 (5897, 5898) Novel Protein sim. GBank gij2132923lpiri S67133 - probable membrane protein YOR240w - yeast (Saccharomyces cerevislae)	95086870 (5899, 5900) Novel Protein sim. GBank gil466102[splP34629]YOJ6_CAEEL - PUTATIVE AMINOPEPTIDASE ZK353,6 IN CHROMOSOME III	87392357 (5901, 5902) Novel Protein sim. GBank gil4688902 emb CAB41450.1
946   94317315 (5891, 589	2947 87362852 (5883, 589	2948 87626527 (5895, 589	2949 88175545 (5897, 586	2850 95086870 (5899, 59	2951 87392357 (5901, 59

2952	95329952 (5903, 5904)	95329952 (5903, 5904) Novel Protein sim. GBank gi 5596693 emb CAB51405.1] -	Contains protein domain (PF00650) - transcriptfactor	ranscriptfactor	264687, 52645156, 21906766, 21906769,
		(ALOSOSO) nypometical protein (nomo sapiens)	Create Constant		264259, 264693, 29331822, 18108365,
					29331823, 00452283, 33837109, 19108508, 29331827, 35696052, 27486262, 284508.
					264905, 20281149, 264908, 264907,
					29331830, 264908, 264909, 35695855,
					264511, 265UU8, 265UU9, 26451U, 264635,   264638 60432220 264838 60433358
				•	264639, 264758, 87168518, 265017.
					22279000, 22279002, 264760, 264563.
					264482, 18108351, 264448, 264288
2953	•	88093575 (5905, 5906) Novel Protein sim. GBank	Contains protein domain (PF00266) - UNCLASSIFIED	UNCLASSIFIED	18108398, 56994075, 22278996, 29331822,
		gil119522 sp P10658 SERC_RABIT - PROBABLE	Aminotransferases class-V		29331824, 29331825, 29331827, 35696052,
		PHOSPHOSERINE AMINOTRANSFERASE (PSAT)			264508, 264905, 264906, 264507, 264510.
		(ENDOMETRIAL PROCESTERONE-INDOCED PROTEIN)			264758, 52646317, 21906754, 33657084,
					52644296, 87168559, 264600, 264760,
					264681, 18108351, 264764, 264369, 264288,
					264687, 21906765, 21906766, 21906767.
					21906768, 21906769, 35695917, 33657023,
					18108364, 52645129, 33657109, 33657349,
					18108374, 263978, 35695855, 264637,
					264638, 87168518, 264482, 264563, 264565
2854	-	88086288 (5907, 5908) Novel Protein sim. GBank	Contains protein domain (PF00019) - 1gf	161	29331822
		gi 4885261 ref NP_005251.1 pGDF9 - growth differentiation	Transforming growth factor beta like		
		factor 9	domain		
2955		87698426 (5909, 5910) Novel Protein sim. GBank gij3452473 (AF084205) -	Contains protein domain (PF00069) - kinase	kinase	264259, 29331822, 29331824, 29331825,
		serine/threonline protein kinase TAO1 [Rattus norvegicus]	Eukaryotic protein kinase domain		29331826, 35696052, 264908, 52644045,
					264512, 60432229, 265018, 265019,
_					55811150, 264769, 21906767, 21906768,
					21906769, 265021, 60170615, 55810764.
2058	_	85780745 (5011 6012) Namel Destrie des CBank	Cathian acolois domois (BEODYB7)		22278006 264250 20111827 264908
}			PX domain		21906768
	_	nexin 7 (Homo sapiens)			
2957		90933301 (5913, 5914) Novel Protein sim. GBank		cadherin	22278999, 264259, 29331824, 29331827.
		gll4503023/reflNP_000089.1 pCPT2 - carnitine			265008, 264595, 264758, 265010, 265011.
		palmitoyltransferase II precursor			264448, 264763, 264683, 264288, 264685,
					18108357, 29148629, 264690, 18108362,
					264693, 18108370, 60431528, 18108374,
					264634, 18108381, 56182323, 18108382,
					18108385, 18108388, 56526486, 87168518,
206	1	to account of the state of the			204402, 204401
0027	_	o/440014 (3913, 3910) NOVel Pfotein sim. Gbank gij424025/jdpj BAA/4907.1  -  /AB0206911 KIAA0884 profein [Homo saniens]			264535, 264567 264631, 264567
		Company of the control of the contro			

263994, 22278997, 264259, 60432049, 29331826, 29331828, 35696052, 29331830, 66712502, 56182435, 265006, 264512, 265008, 265008, 265009, 60433358, 60433438, 264595, 265017, 265018, 264683, 264288, 264769, 21906766, 21906767, 21906769, 265020, 60170615, 264692, 27486265, 18108374, 65274791, 35695855, 83373044, 56526486, 60432113	35696286, 56182435, 87168474, 265010, 60170615, 35696423, 56182323, 18108383, 87168518, 264483	22278997, 22278999, 264259, 29331822, 29331824, 29331826, 29331828, 264907, 264908, 5264908, 33657402, 21908754, 87188474, 265011, 87168559, 265017, 21908769, 265020, 60170615, 264692, 33557023, 35695763, 18108370, 181083385, 87168518, 22279002, 264564, 264567	35696286, 22278997, 264092, 264094, 264259, 29331824, 66714117, 29331825, 60432289, 29331826, 29331827, 29331825, 35896052, 264508, 264508, 264509, 264512, 264593, 264594, 60433438, 264510, 264597, 264593, 264602, 264634, 264288, 264766, 264691, 264569, 264634, 264638, 264634, 264638, 264634, 264638, 264634, 264565, 264567, 264565, 264567, 264563, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 264564, 26464, 26464, 26464, 26464, 26464, 26464, 26464, 26464,	18108392, 56994075, 22278998, 22278999, 29331827, 29331822, 29331822, 29331827, 29331828, 265007, 265008, 264592, 264594, 21906754, 265018, 264760, 264687, 29148627, 29148784, 265020, 33657023, 264693, 65274620, 33657182, 27486261, 264629, 55810764, 35696423, 264555, 264637, 264637, 264557, 264558, 264563	264259, 29331828, 33657402, 265017, 265018, 264692, 18108368, 35698423, 83373044, 18108388
	•		UNCLASSIFIED 2 2 3 3 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2		UNCLASSIFIED
Contains protein domain (PF00566) - TBC domain		Contains protein domain (PF00400) - Itranscriptfactor WD domain, G-bela repeat		Contains protein domain (PF00010) - Iranscriptfactor Helix-loop-helix DNA-binding domain	
2959   95109420 (5917, 5918) Novel Protein sim. GBank gilg88221 (U33005) - Tbc1 (Mus   Contains protein domain (PF00566) - loncogene musculus)		95413416 (5921, 5922) Novel Protein sim. GBank gil5596646 emb CA805177.2  - Contains protein domain (P (282266) predicted using Genefinder; similar to WD domain, WD domain, G-beta repeat G-beta repeats [Caenorhabdilis elegans]		95313464 (5925, 5926) Novel Protein sim. GBank gil4240223 dbj BAA74890.1  - (AB020674) KIAA0867 protein [Homo saplens]	
95109420 (5817, 5918)		95413416 (5921, 5922 <u>)</u>		85313464 (5925, 5926)	94324617 (5927, 5928)
2959	2960	2961	2962	2963	2964

2965	80384762 (5929, 5930)			transcriptfactor	264259, 29331826, 264508, 264509, 264905, 264907, 264908, 264909, 264511, 265008, 264910, 264591, 264593, 264594, 33657402, 265011, 264760, 264762, 264764, 264288, 264685, 264766, 264692, 33657109, 264628, 264639, 264631, 264632, 264639, 264631, 264638, 264639, 264638, 264639,
	91725248 (5931, 5932)				60432289, 264682, 264448
2967	94658303 (5933, 5934)	94658303 (5933, 5934) Novel Protein sim. GBank gi 624225 (U19181) - Rabin3 [Rattus norvegicus]		UNCLASSIFIED	264488, 264508, 264509, 264908, 264909, 264511, 264910, 264594, 264768, 85658542, 264762, 264764, 265021, 264556, 18108381, 264564, 264488
2968	95302776 (5935, 5936)	I/AF15188 - (AF151881) CGI-123	Contains protein domain (PF00097) Inc finger, C3HC4 type (RING finger)		264687, 52645156, 21908765, 52646365, 21906767, 18108398, 35696423, 22278996, 3569628, 22278999, 3569628, 22278999, 265021, 265022, 264093, 264093, 264698, 2644150, 264259, 33657023, 2545080, 264693, 29331825, 33109954, 52645129, 29331826, 21906754, 33657182, 29331827, 29331826, 21906754, 33657182, 29331827, 29331826, 21906754, 33657182, 29331827, 25238005, 27486262, 87168518, 87168474, 265019, 87168559, 265019, 22278000, 265019, 22278000, 265019, 264448, 66712502, 264568, 264369, 264288
		95310957 (5937, 5938) Novel Protein sim. GBank gij3024743 sp O24734 THSA_SULS7 - THERMOSOME, ALPHA SUBUNIT (CHAPERONIN ALPHA SUBUNIT)		- da	52646842, 22278996, 22278998, 22278999, 60432049, 264259, 29331824, 29331825, 29331826, 29331824, 29331825, 29331826, 29331826, 29331826, 29331826, 29331826, 264099, 2644046, 264091, 26448, 26498, 264359, 2644229, 21906766, 21906768, 21906769, 29148784, 265021, 52644150, 264691, 33657109, 18108374, 56182323, 60170394, 87168518, 60432113, 22279000
2970	88088071 (5939, 5940)	88088071 (5939, 5940) Novel Protein sim. GBank gij3165407 (AC004755) - [6s37502_1 [Homo sapiens]	Contains protein domain (PF00046) - homeobox Homeobox domain	нотеорох	

-	94196930 (5941, 5942)	2971 94196930 (5941, 5942) Novel Protein sim. GBank gil728837 sp P39194 ALU7_HUMAN - IIII ALU SUBFAMILY SQ WARNING ENTRY IIII		(m)	264488, 56182575, 35696286, 56994075. 29331824, 29331826, 29146499, 284508. 264905, 264907, 264112, 264910, 21906754. 87168559, 265018, 265019, 18108351. 264689, 21906765, 21906767, 21906768.
				14 14 W	265020, 265021, 60170615, 18106364, 264628, 264629, 18108374, 264636, 264556, 264558, 83373044, 18108384, 18108385, 87168518, 264564, 264567
2972	86625943 (5943, 5944)	86625943 (5943, 5944) Novel Protein sim. GBank gif728836spP39193JALU6_HUMAN - IIII ALU SUBFAMILY SP WARNING ENTRY IIII	W.		265017, 35695917, 265021, 33657109, 22279002, 264563
2973	91215301 (5945, 5946)	91215301 (5945, 5946) Novel Protein sim. GBank gij2746789 (AF040642) - No definition line found (Caenorhabditis elegans)	<b>5</b>		29331822, 264692, 33657349, 55811576, 264563
2974		91673002 (5947, 5948) Novel Protein sim. GBank gil786117 (L41834) - nuclear protein [Ensis minor]		UNCLASSIFIED	18108391, 52644501, 50182575, 50181600, 22278995, 22278996, 35696286, 22278997, 22278996, 25278996, 22278997, 22278998, 22278999, 264259, 52645080, 29331824, 29331825, 66714117, 60424269, 26331826, 26331827, 26331828, 3569652, 266712502, 264908, 52644045, 265007, 265009, 60433438, 33109954, 21906754, 55811386, 52644296, 87168474, 87168559, 265017, 265018, 265019, 18108351, 264448, 264369, 265019, 264622, 1906765, 21906767, 21906768, 35695917, 265020, 265021, 52644150, 264691, 264692, 33657023, 27486262, 2748624, 35695763, 18108370, 18108376, 55810764, 55811576, 35696423, 35695855, 264535, 264587, 356964332, 264581, 364482,
2975	95325213 (5949, 5950)	95325213 (5949, 5950) Novel Protein sim. GBank gij3880812 emb CAA19508  - Contains protein domain (AL023839) similar to HECT-domain (ubiquitin-transferase).; HECT-domain (ubiquitin-cDNA EST yk880410.5 comes from this gene	Contains protein domain (PF00632) - ubiquitin HECT-domain (ubiquitin- transferase).		29331824, 29331827, 29331828, 264910, 85658542, 265011, 265018, 264448, 264288, 264769, 21908767, 265020, 264691, 264559, 83373044
2976		87771202 (5951, 5952) Novel Protein sim. GBank gil5679136 gb AAD46874.1 AF16093 - (AF160934) BcDNA.LD14189 [Drosophila melanogaster]		transport	22278996, 264906, 265007, 265010, 265011, 265017, 265017, 265019, 18108351, 264685, 264689, 18108370, 264639, 18108385
2977	91725254 (5953, 5954)	91725254 (5953, 5954) Novel Protein sim. GBank gij5262751jemb CAB45690.1  - (AJ243177) Xenopus RPA interacting protein alpha [Xenopus laevis]		UNCLASSIFIED	264509, 264288

2978	87332059 (5955, 5956)	2978   87332059 (5955, 5956)   Novel Protein sim. GBank gil746549 (U23522) - No	Contains protein domain (PF00480) - UNCLASSIFIED		22278995, 22278996, 22278997, 22278999, 242759, 6043228, 29331827, 29146499
					56182435, 265006, 265007, 265009,
					60433356, 60433438, 21906754, 265010,
					265011, 265017, 265018, 265019, 264288,
					264685, 264688, 21906765, 21906768.
					21906767, 21906768, 21906769, 265020,
					265021, 265022, 35696423, 264639,
					60432113, 22279000, 22279002
2979	91725256 (5957, 5958)	91725256 (5957, 5958) Novel Protein sim. GBank gij5262751 emb CAB45690.1  -		complement	264488, 65274572, 56994075, 22278999,
					264093, 29331822, 29331824, 264288,
_		[Xenopus laevis]			55811957, 33657023, 33657109, 18108370,
					55811576, 56182323, 60432113, 264482
2980	86296600 (5959, 5960)				265009, 21906767, 263981, 22279000
2981	-			UNCLASSIFIED	264629, 264564
2982		95303675 (5963, 5964) Novel Protein slm. GBank			22278995, 56994075, 22278996, 22278997.
		gil4929767[qb]AAD34144.1 AF15190 - (AF151907) CGI-149	6		22278998, 22278999, 264092, 29331824,
		protein [Homo sapiens]			29331827, 29331828, 264905, 264591,
					264592, 264594, 264595, 264596, 33657084.
					264448, 21906765, 21906766, 21906767.
		•			21906768, 21906769, 265020, 265022,
					18108365, 33657182, 33657349, 35696423,
					83373044, 22279000, 22279002
2983	91725258 (5965, 5966)	91725258 (5965, 5966) Novel Protein sim. GBank gi[5262751 emb[CAB45690.1] -			60424179, 52646842, 18108398, 22278997,
		(AJ243177) Xenopus RPA Interacting protein alpha			264093, 60432049, 264259, 29331822,
		(Xenopus laevis)			60432289, 33656970, 264905, 52644045,
					265006, 60431735, 87168474, 265018.
					265019, 18108351, 264448, 21906765.
					21906768, 35695917, 33657023, 52645129,
					18108370, 35696423, 83373044, 56526486.
_					60432113, 264404, 22279002
2984				ATPase_associated	
		to C. elegans F11A10.5; 80% similarity to Z68297			
900		Consider the control of the control		Ī	
2882		Novel Protein sim. GBank gi[103160 pir  522126 - finger		UNCLASSIFIED	264910, 55812038, 56181562, 55811957.
		(protein unkempt - iruit ily (Urosopnila melanogaster)		•	204628, 53810/04, 204632, 204633, 60432113
2986					55811957, 264566
7987				ACCICION I	264360
2				UNCLASSIFIED	264369

264488, 65274572, 22278995, 22278996, 22278997, 22278999, 264092, 264094, 264259, 60432049, 26331824, 29331826, 60432289, 35696052, 28331824, 29331826, 60432289, 35696052, 28331828, 264107, 264905, 264907, 265008, 66712502, 264828, 264905, 56182435, 265008, 265007, 265008, 260170831, 60433229, 2484593, 60433356, 265011, 87168559, 265017, 265018, 264682, 264448, 264369, 265017, 265021, 265031, 2646937, 264635, 264635, 264635, 264565, 264565, 264567, 264566, 264567, 264566, 264567, 264566, 264567, 264566, 264567, 2	22278996, 22278997, 264905, 264511, 60170831, 264593, 265019, 21906765, 21906767, 21906768, 18108374	265007, 264512, 18108351, 264288, 264689, 265020, 264691, 33657023, 33657109	264563	264259, 265019, 264689, 18108385	264488, 29331822, 265017, 264761, 21906769, 65274791, 261981, 264565	22278994, 22278995, 56994075, 22278997, 22278999, 284259, 29331822, 29331824, 29331825, 29331825, 29331828	265006, 265009, 264910, 33109954, 87168474, 87168559, 265018, 285019.	264448, 264288, 21906766, 21906767, 21906768, 21906768, 263021, 265022, 33657023, 264693, 35695855, 83373344	18108385, 22279000, 264565, 264566 284905, 264907, 265019, 18108351, 264683	65274572, 35696286, 264259, 29331824,	35696052, 29146499, 264508, 264907, 265007, 265008, 60433438, 18108348.	265017, 264681, 264683, 264288, 264766,	264769, 264689, 35695917, 60170615, 33657023, 264692, 264634, 264555	18108381, 18108382, 18108388, 264484
UNCLASSIFIED	kinase	опсодепе	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	transport				helicase				
.i.		Contains protein domain (PF00071) - oncogene Ras family									DEAD/DEAH box helicase			
2988 95357753 (5975, 5976) Novel Protein sim. GBank gil4679028 gb AAD27002.1  - (AF077207) HSPC021 [Homo sapiens]	91225118 (5977, 5978) Novel Protein sim. GBank gi 113671 sp P23964 ALUF_HUMAN - IIII ALU CLASS F WARNING ENTRY IIII	87330444 (5979, 5980) Novel Protein sim. GBank gi 2829836 sp P97348 RHOD_MOUSE - RHO-RELATED GTP-BINDING PROTEIN RHOD				94136634 (5987, 5988) Novel Protein sim. GBank 91 2496549 sp Q50658 YU02_MYCTU - HYPOTHETICAL 29.7 KD PROTEIN CY339.02			87591070 (5989, 5990) Novel Protein sim. GBank gil2734081 (AF000195) - similar	91013798 (5991, 5992) Novel Protein sim. GBank gij2829912 (AC002291) - Similar	A I P-dependent KNA Helicase [Arabidopsis thaliana]			
95357753 (5975, 5976)	91225118 (5977, 5978)	87330444 (5979, 5980)	94325361 (5981, 5982)	_		9,4136634 (5987, 5988)					-			
2988	5883	2890	2002	7000	2882	2894			2985	2996				

2007	R7627440 (5003 5004)	Monet Dentain ein CBank allasabs Statistaans saat 11.	homeohox	264488 56182575 284259 66714117
		(AB023221) KiAA1004 protein IHomo saplensi		29331828, 35696052, 264508, 264509,
				264907, 264908, 265006, 87168474, 265019,
				264448, 264682, 264685, 264766, 21906764,
				21906766, 21906768, 21906769, 27486261,
				18108374, 35696423, 264634, 264635,
				264636, 264557, 18108385, 87168518
2998	88095381 (5995, 5996)	88095381 (5995, 5996) Novel Protein sim. GBank gil3947589 emb CAA22252  -	UNCLASSIFIED	52646365, 22278997, 264508, 264906,
		(AL034364) cDNA EST yk255b9.3 comes from this gene;		18108351, 21906765, 21906767, 18108370,
		cDNA EST yk255b9.5 comes from this gene; cDNA EST		18108374, 35696423, 264636, 264639
		EMBL:M75923 comes from this gene [Caenorhabditis		·
		(elegans)		
2999		Novel Protein slm. GBank	UNCLASSIFIED	56182575, 22278996, 29147620, 29331825,
		gij115408jspjP18835jcC19_CAEEL - CUTICLE		29146498, 29146499, 264905, 66712502,
		COLLAGEN 19		265006, 265009, 21906754, 85658542,
		•		18108351, 29148627, 29148629, 60170615,
			•	33657109, 27486262, 18108370, 18108374,
				264556, 264557, 264558, 60170394,
				18108385, 264563
3000		95099370 (5999, 6000)  Novel Protein sim. GBank gi 1163174 (U32575) - similar to	UNCLASSIFIED	264887, 22278997, 22278999, 264259,
		yeast Sec6p, Swiss-Prot Accession Number P32844; similar		29331822, 29331824, 35696052, 29146498,
		to mammalian B94, Swiss-Prot Accession Number Q03169;		264508, 264905, 264906, 264907, 264908,
		Method: conceptual translation supplied by author (Rattus		264909, 264510, 264511, 265006, 265007,
		novegicus]		265008, 265009, 264810, 33657402, 264757,
				264595, 264596, 264758, 21906754, 265011,
				264600, 265017, 265018, 264605, 265019.
_				264760, 264761, 264762, 264681, 264682,
				264764, 264288, 264685, 264766, 264686,
				264768, 264769, 21906765, 21906768.
		•		35695917, 265020, 264691, 264692,
				33657023, 264693, 33657109, 33657182.
				27486261, 264628, 264629, 18108374,
				18108376, 35696423, 35695855, 264630,
			-	264631, 264632, 264634, 264635, 264636,
				264637, 264558, 284638, 264639, 60170394,
				83373044, 20798451, 22279002, 264553,
	- 1			264486, 264567
300 100		88078454 (6001, 6002) Novel Protein sim. GBank gi 2078470 (AC002073) -	cathepsin	18108394, 52646842, 56182575, 29331824.
_		Putative gene. Genscan predictions confirmed by EST		29331825, 29331827, 264910, 33109954,
		splicing.; coded for by human cDNAs AA122029		52644296, 265017, 265019, 264288, 265020,
		(NID:g1678048), D31562 (NID:g644442), AA158721		265021, 52644150, 284692, 35695763,
		(NID:91733515), R59640 (NID:9830335) and F13082		55810764, 35696423, 56182323, 18108387.
		(NID:g709111) [Homo sapiens]		264563, 264564
3002	87718167 (6003, 6004)	87718167 (6003, 6004) Novel Protein sim. GBank gij3599478 (AF085185) - Myosin-	UNCLASSIFIED	264488, 29331824, 29331825, 29331826,
		IA [Acanthamoeba castellanii]		29331827, 29331828, 264906, 264510,
				265009, 21906754, 264682, 264688.
				133657023, 264565

	29331830, 21906769, 264691, 33657109, 263972, 18108385		23) - MHC 264259, 28331822, 264512, 21906754, 265018, 284687, 21906765, 264691, 264555, 264558, 18108385		264905, 264593, 264766, 264636				1607704, 607001
Contains protein domain (PPU 391) - cottagen Collagen triple helix repeat (20 copies)		Contains protein domain (PF01360) - oxygenase Monooxygenase	Contains protein domain (PF00023) - MHC Ank repeat	Contains protein domain (PF00096) - Iranscriptfactor Zinc finger, C2H2 lype		Contains protein domain (PF0050 AMP-binding enzyme	Contains protein domain (PF01923) - UNCLASSIFIED Protein of unknown function	Contains protein domain (PF00153) - transport Mitochondrial carrier proteins	
1 / 54 55 5 (0.30 25 2) - Collagen musculus]	88066876 (6007, 6008) Novel Protein sim. GBank gi 2224629 db  BAA20802  - (AB06876 (6007, 6008)	(AF132944) CGI-10	87422224 (6011, 6012) Novel Protein sim. GBank gij3930525 (AF064447) - sex- determination protein homolog Fem1a (Mus musculus)	90936005 (6013, 6014) Novel Protein sim. GBank gil2565052 (U80738) - CAGH1a [Homo saplens]		91213387 (6017, 6018) Novel Protein sim. GBank gil3127193 (AF062389) - kidney- Contains protein domain (PF00501) - synthase specific protein [Rattus norvegicus]  AMP-binding enzyme	95317217 (6019, 6020) Novel Protein sim. GBank gil4927370 gb AAD33084.1 AF06797 - (AF067972) DNA Cytosine methylkransferase 3 alpha (Homo sapiens)	94323597 (6021, 6022) Novel Protein sim. GBank gil5052319 gblAAD38501.1 AF11883 - (AF118838) citin; adult-onset type II citrullinemia protein [Homo sapiens]	
88848078 (8003, 8008)	88066876 (6007, 6008)	87794843 (6009, 6010)	87422224 (6011, 6012)	90936005 (6013, 6014)	80416249 (6015, 6016)	91213387 (6017, 6018)		94323597 (6021, 6022)	
3003	3004	3005		3007	3008	3006	3010	3011	

264488, 263994, 35696286, 22278997, 264259, 29331824, 60424269, 66714117, 35698052, 284906, 264906, 264907, 264908, 264909, 56182435, 264511, 244512, 264910, 264591, 264591, 264591, 264591, 264591, 264591, 264591, 264591, 264592, 264592, 265010, 265018, 264004, 264760, 264682, 264683, 264686, 264689, 264628, 264629, 264631, 264632, 264639, 264631, 264631, 264634, 264636, 264636, 264631, 264631, 264638, 264636, 264636, 264565, 264567, 264631, 264638, 264636, 264636, 264565, 264636, 264565, 264567, 264655, 264636, 264565, 264567, 264565, 264565, 264567, 264565, 264565, 264567, 264565, 264567, 264565, 264567, 264565, 264567, 264565, 264567, 264565, 264567, 264565, 264567, 264565, 264567, 264565, 264567, 264565, 264567, 264565, 264567, 264565, 264567, 264567, 264565, 264567, 26457, 2		22278995, 22278996, 22278997, 264259, 29331824, 29331828, 264906, 265007, 265008, 264910, 265011, 265017, 265019, 264691, 33657109, 18108370, 35695855, 264556, 264564	52644507, 52646842, 56994075, 52645080, 29331822, 29331824, 35696052, 33656970, 52644045, 264596, 33657084, 265017, 265019, 52644229, 21906767, 35695917, 52644150, 33657023, 33657109, 27486261, 27486262, 27486265, 35695763, 35695763, 35695763, 35695763, 36957635, 87168518	264488, 18108397, 22278996, 35696286, 22278999, 264259, 29331822, 60432289, 264908, 29331832, 60432289, 264908, 29331830, 264908, 56182435, 265006, 265007, 265008, 265009, 264591, 60433436, 626011, 8716859, 265017, 265018, 265011, 8716859, 265020, 265020, 21906766, 21906766, 21906769, 265020,
Iranscriptfactor	ATPase_associaled		UNCLASSIFIED	·
Contains protein domain (PF00400) - Iranscriptfactor				
433374_1 [Homo sapiens]	79877263 (6027, 6028) Novel Protein slm. GBank gij3878374 emb CAA93081 - (Z68879) Similarity to Yeast Chi12p protein (PIR Acc. No. S54453); cDNA EST EMBL:D27950 comes from this gene; cDNA EST EMBL:D27949 comes from this gene; cDNA EST EMBL:D33447 comes from this gene; cDNA EST EMBL:D33316 comes from this gene; cDNA EST		87759945 (6031, 6032) Novel Protein sim. GBank gij1168819jspiP41733jCC91_YEAST - CELL DIVISION CONTROL PROTEIN 91	95011154 (6033, 6034) Novel Protein sim. GBank gil4589658 dbj BAA76851.1  - (AB023224) KIAA1007 protein [Homo sapiens]
3013   91238799 (6025, 6026)   Novel Protein sim. GBank gi R33374_1 [Homo saplens]	3014 79877263 (6027, 6028) N (6) (7) (6) (7) (7) (7) (7) (7) (7) (7) (7) (7) (7	86995466 (6029, 6030)	3016   67758945 (6031, 6032)   677589450 (6031, 6032)   67758945 (6031, 6032)   67758945 (6031, 6032)	3017 95011154 (6033, 6034) h

	2040	11072801 (6025 6026)				264558
gij3414809 (AF061529) - rjs [Mus Contains protein domain (PF00415) - ATPase_associated gij3414809 (AF061529) - rjs [Mus Contains protein domain (PF00415) - ATPase_associated condensation (RCC1)  igj3880889[emb]CA809005] - Gondensation (RCC1)  igj3880889[emb]CA809005] - Gondensation (RCC1)  igj28601 (L1275) - selected as a full and it it is gene (Candans protein domain (PF00614) - UNCLASSIFIED (Contains protein domain (PF00614) - UNCLASSIFIED (Contains protein domain (ACAO of DNA and it is it is add it it it is add it it is add it it it is add it it it it is add it	: I:	073691 (0035, 0030)				64560 K2644507 18108304 65274572
gij3414809 (AF061529) - rjs [Mus Contains protein domain (PF00415) - ATPase_associated Regulator of chromosome condensation (RCC1)  gij3880899jemb[CAB09005] - 6045 contains protein domain (PF00415) - ATPase_associated gij3880899jemb[CAB09005] - 6045 contains protein domain (PF00614) - UNCLASSIFIED (Contains domain (PF00614) - UNCLASSIFIED (	\$	148231 (6037, 6038)		2		04503, 36044301, 10100334, 03414374,
gij3414809 (AF061529) - fjs [Mus Contains protein domain (PF00415) - ATPase_associated Regulator of chromosome condensation (RCC1) [3880889[emb[CA8038005] - 684.5 comes from this gene; cDNA EST mis gene; cDNA EST in gene [Caenorhabditis elegans] [4] YEAST - A-AGGLUTININ Phospholipase D. Active site motif PRECURSOR [5] PRECURSOR [6] 1295671 (L11275) - selected as utant of the subunit Ac46 of DNA se 1 and III [Saccharomyces			Unknown gene product (Homo sapiens)			61825/5, 222/6994, 222/0995, 509940/5,
gij3414809 (AF061529) - rjs [Mus Contains protein domain (PF00415) - ATPase_associated Regulator of chromosome condensation (RCC1) gij3880889[emb[CA609005] - 664.5 comes from this gene; cDNA m this gene; cDNA EST his gene (Caenorthabdilis elegans) 1.1 YEAST - A-AGGLUTININ Phospholipase D. Active site motif PRECURSOR gij295671 (L11275) - selected as utant of the subunit AC40 of DNA se I and III [Saccharomyces					<del>-</del>	2278998, 22278999, 264259, 29331822,
gij3414809 (AF061529) - rjs [Mus Contains protein domain (PF00415) - ATPase_associated Regulator of chromosome condensation (RCC1) gij3880889[embjCA809005] - Gones from this gene: cDNA m this gene: cDNA m this gene: cDNA EST his gene: cDNA EST his gene: cDNA EST his gene [Caenorhabditis elegans] A1_YEAST - A-AGGLUTININ Phospholipase D. Active site motif PRECURSOR Gij295671 (L11275) - selected as utant of the subunit AC40 of DNA se I and III [Saccharomyces				-	<u> </u>	9331824, 60432289, 29331827, 264908,
gij3414809 (AF061529) - rjs [Mus Contains protein domain (PF00415) - ATPase_associaled Regulator of chromosome condensation (RCC1)  gij3880889[emb]CAB09005] - Gondensation (RCC1)  comes from this gene; cDNA EST his gene; cDNA EST his gene; cDNA EST his gene; cDNA EST his gene [Caenorhabdilis etegans]  Contains protein domain (PF00614) - UNCLASSIFIED PRECURSOR  Gij295671 (L11275) - selected as utant of the subunit AC40 of DNA as 1 and III [Saccharomyces se 1 and III [Saccharomyces						6182435, 265007, 265009, 60432229,
gij3414809 (AF061529) - rjs [Mus Contains protein domain (PF00415) - ATPase_associated Regulator of chromosome condensation (RCC1)  Sequiator of chromosome condensation (RCC1)  Sequiator of chromosome condensation (RCC1)  This gene (Canom this gene; DNA m this gene; CDNA EST his gene [Caenorhabditis elegans]  At_YEAST - AAGGLUTININ Phospholipase D. Active site motif PRECURSOR  Signatory (L11275) - selected as utant of the subunit AC40 of DNA as 1 and III [Saccharomyces as 1 and III [Saccharomyces			•			64593, 60433356, 55812038, 21906754,
gij3414809 (AF061529) - rjs [Mus Contains protein domain (PF00415) - ATPase_associated Regulator of chromosome condensation (RCC1)  8gij3880889[emb]CAB09005] - 664.5 comes from this gene: connes from this g						7168474, 265011, 87168559, 265017,
gij3414809 (AF061529) - /js [Mus Contains protein domain (PF00415) - ATPase_associated Regulator of chromosome condensation (RCC1)  640.5 comes from this gene; cDNA m this gene; cDNA EST his gene [Caenorhabditis etegans]  1. YEAST - A-AGGLUTININ Phospholipase D. Active site motif PRECURSOR  1. Gontains protein domain (PF00614) - UNCLASSIFIED PRECURSOR  1. YEAST - A-AGGLUTININ Phospholipase D. Active site motif PRECURSOR  1. Selectied as utant it [Saccharomyces as I and Itt [Saccharomyces]				-		65018, 265019, 264681, 18108351, 264448,
gij3414809 (AF061529) - 1/s [Mus Contains protein domain (PF00415) - AFPase_associated Regulator of chromosome condensation (RCC1)  1913880889 emb CAB09005  - 664.5 comes from this gene; cDNA EST in this gene [Caenombaditis elegans]  1.1 YEAST - A-AGGLUTININ Phospholipase D. Active site motif PRECURSOR  1.2 YEAST - A-AGGLUTININ Phospholipase D. Active site motif PRECURSOR are in and it [Saccharomyces]						64682, 264683, 18108354, 264685, 264687,
gij3414809 (AF061529) - 1js [Mus Contains protein domain (PF00415) - ATPase_associated Regulator of chromosome condensation (RCC1)  gij3880889jemb[CAB09005] - 664.5 comes from this gene; cDNA EST mits gene; cDNA EST in this gene; cDNA EST in this gene [Caenorhabditis elegans]  Contains protein domain (PF00614) - UNCLASSIFIED PRECURSOR  PRECURSOR  UNCLASSIFIED OINCLASSIFIED PRECURSOR  gij293671 (L11275) - selected as utant of the subunit AC40 of DNA as 1 and Itil [Saccharomyces						64689, 21906766, 21906768, 21906769,
gij3880889jemb[CAB09005] - 64.5 comes from this gene; condensation (RCC1)  gij3880889jemb[CAB09005] - 64.5 comes from this gene; connes from this gene; connes from this gene; cDNA  m this gene; cDNA EST  his gene [Caenorhabditis elegans]  Ly YEAST - A-AGGLUTININ  PRECURSOR  gij295671 (L11275) - selected as utant of the subunit AC40 of DNA  se I and III [Saccharomyces						52644150 264690 264691 33657023.
gij3810889jemb[CAB09005] - rjs [Mus Contains protein domain (PF00415) - ATPase_associated Regulator of chromosome condensation (RCC1) gij3890889jemb[CAB09005] - Gondensation (RCC1) gij380889jemb[CAB09005] - Gondensation (RCC1) mis gene; cDNA EST his gene; cDNA EST his gene [Caenorhabditis elegans] v1_YEAST - A-AGGLUTININ Phospholipase D. Active site motif PRECURSOR gij295871 (L11275) - selected as utant of the subunit AC40 of DNA SI PRECURSOR						84692 264693 33657109 52645129.
gij38989lembjCAB09005j - rjs [Mus Contains protein domain (PF00415) - ATPase_associated Regulator of chromosome condensation (RCC1) gij388089jembjCAB09005j - 664.5 comes from this gene; comes from this gene; comes from this gene; comes from this gene; comes from this gene [Caenorhabditis elegans] is gene [Caenorhabditis elegans] his gene [Caenorhabditis elegans]  Contains protein domain (PF00614) - UNCLASSIFIED PRECURSOR  INCLASSIFIED UNCLASSIFIED UNCLASSIFIED (Contains protein domain (PF00614) - UNCLASSIFIED (Caenorhabditis elegans) are all motif precursors.						13657340 364630 65274701 264634
gij3414809 (AF061529) - rjs [Mus Contains protein domain (PF00415) - ATPase_associated Regulator of chromosome condensation (RCC1)  gij3880889[emb]CAB09005] - Gondensation (RCC1)  Gones from this gene; cDNA m this gene; cDNA EST his gene [Caenorhabditis elegans]  his gene [Caenorhabditis elegans]  At _YEAST - A-AGGLUTININ Phospholipase D. Active site motif PRECURSOR  gij295671 (L11275) - selected as utant of the subunit AC40 of DNA as I and till [Saccharomyces						33031343, E04023, CJE 14731, E04034,
gij3414809 (AF061529) - rjs [Mus Contains protein domain (PF00415) - ATPase_associated Regulator of chromosome condensation (RCC1)  gij3880889 emb CAB09005  - Gondensation (RCC1)  m this gene; cDNA EST his gene; cDNA EST his gene [Caenorhabditis elegans]  t. YEAST - A-AGGLUTININ Phospholipase D. Active site motif PRECURSOR  gij295671 (L11275) - selected as utant of the subunit AC40 of DNA  se I and III [Saccharomyces						22244332, 30102323, 10100303, 07100310, 022320, 032320, 03232000, 032320000, 064463
gij3880889jembjCAB09005j - Gondensation (RCC1)  864.5 comes from this gene; cDNA m this gene; cDNA EST his gene; cDNA gij295671 (L11275) - selected as utant of the subunit AC40 of DNA se i and III [Saccharomyces		4210261 (E020 E040)	Mana Pentale sim CBank all 2414800 (AE061620) - ris (Mus.	Contains protein domain (PE00415) -	ATPace associated	P64488 283994 35896288 264259 264508.
gij3860889jembjCAB09005j - 664.5 comes from this gene; cDNA m this gene; cDNA EST his gene [Caenomabditis elegans] his gene [Caenomabditis elegans]  1_YEAST - A-AGGLUTININ PRECURSOR gij295671 (L11275) - selected as utant of the subunit AC40 of DNA  19		7310531 (0033, 0040)	misculies	Regulator of chromosome		264905, 264509, 264906, 264907, 264908,
gij3880889jembjCAB09005j - 604.5 comes from this gene; diglascontable and the subunit AC40 of DNA gij295671 (L11275) - selected as gij295671 (L1275) - selected as gij295671 (L11275) - selected as gij29571 (L11275) - se				Condensation (RCC1)		264909, 264510, 264910, 60174639, 264600,
gij3880889jembjCAB09005j - 604.5 comes from this gene; dignary of the subunit action to the subunit Actor of DNA gij295671 (L11275) - selected as subunit Actor of DNA gij295671 (L11275) - selected as subunit Actor of DNA gij295671 (L11275) - selected as subunit Actor of DNA						264603, 264760, 264762, 264682, 264763.
gij3880889jemb[CAB09005] - 604.5 comes from this gene; comes from						264764, 264288, 264369, 264766, 264687,
gij3880889 emb[CAB09005  - 6d4.5 comes from this gene; cDNA m this gene; cDNA EST his gene [Caenorhabdilis elegans]  vt_YEAST - A-AGGLUTININ PRECURSOR  gij295671 (L11275) - selected as utant of the subunit AC40 of DNA ig and III [Saccharomyces]						264688, 264769, 55811957, 35695917,
gij3880889jembjCA809005j - 6d4.5 comes from this gene; comes from this gene; cons from this gene; contains protein domain (PF00614) - UNCLASSIFIED PRECURSOR gij295671 (L11275) - selected as utant of the subunit AC40 of DNA se f and iii [Saccharomyces						33657023, 264628, 35696423, 35695855,
gij3880889jemb[CAB09005] - 664.5 comes from this gene; comes from						264630, 264632, 264634, 264635, 264636,
gij3880889jemb[CAB09005j - 6d4.5 comes from this gene; contains protein domain (PF00614) - UNCLASSIFIED  Contains protein domain (PF00614) - UNCLASSIFIED  PRECURSOR  gil295671 (L11275) - selected as ulant of the subunit AC40 of DNA ise f and till (Saccharomyces						264637, 264556, 264557, 264638, 264639,
gij3860889jembjCAB09005j - 6d4.5 comes from this gene; comes from						33373044, 18108385, 264564, 264567,
gij3860889jembjCAB09005j - 6d4.5 comes from this gene; comes from this gene; cDNA m this gene; cDNA EST his gene [Caenorhabditis elegans]  LI_YEAST - A-AGGLUTININ PRECURSOR  gij295671 (L11275) - selected as ugat 3 and III [Saccharomyces]						264486
664.5 comes from this gene;  comes from this gene; cDNA  m this gene; cDNA  Contains protein domain (PF00614) - UNCLASSIFIED  PRECURSOR  gil295671 (L11275) - selected as gil295671 (L11275) - selected as gil295671 (L1275) - selected as selected a		80478512 (6041, 6042				264769, 264629, 264482
comes from this gene; cDNA m this gene; cDNA EST his gene [Caenorhabditis elegans]  Contains protein domain (PF00614) - UNCLASSIFIED Contains protein domain (PF00614) - UNCLASSIFIED PRECURSOR gi[295671 (L11275) - selected as gi[295671 (L1275) - selected as to and iii [Saccharomyces]			(Z95559) cDNA EST yk236d4.5 comes from this gene;			
his gene; cDNA EST his gene [Caenorhabditis elegans] Ly YEAST - A-AGGLUTININ PRECURSOR gi[295671 (L11275) - selected as ulant of the subunit AC40 of DNA  This gene [Caenorhabditis elegans]  Contains protein domain (PF00614) - UNCLASSIFIED  UNCLASSIFIED  UNCLASSIFIED  UNCLASSIFIED  se t and III [Saccharomyces			CDNA EST EMBL:C13455 comes from this gene; cDNA			
his gene [Caenorhabdilis elegans]  UNCLASSIFIED  UNCLASSIFIED  Contains protein domain (PF00614) - UNCLASSIFIED  PRECURSOR  gi[295671 (L11275) - selected as utant of the subunit AC40 of DNA se I and III [Saccharomyces			EST yk329g6.5 comes from this gene; cDNA EST			
UNCLASSIFIED Contains protein domain (PF00614) - UNCLASSIFIED PRECURSOR gil295671 (L11275) - selected as utant of the subunit AC40 of DNA se f and III (Saccharomyces			CEMSH45R comes from this gene [Caenorhabditis elegans]			
Contains protein domain (PF00614) - UNCLASSIFIED PRECURSOR gi[295671 (L11275) - selected as ulant of the subunit AC40 of DNA se f and III [Saccharomyces		87718500 (6043 6044			INCIASSIEIED	264259 29331826 29331828 264288
Contains protein domain (PF00614) - UNCLASSIFIED PRECURSOR gi[295671 (L11275) - selected as utant of the subunit AC40 of DNA se I and III [Saccharomyces						264566
PRECURSOR  gi[295671 (L11275) - selected as ulant of the subunit AC40 of DNA se f and III [Saccharomyces		95305484 (6045, 6046	Novel Protein sim. GBank	Contains protein domain (PF00614) -	UNCLASSIFIED	264488, 22278995, 35696286, 22278997,
PRECURSOR  UNCLASSIFIED gi[295671 (L11275) - selected as ulant of the subunit AC40 of DNA se i and iii (Saccharomyces			gil416592lsplP32323JAGA1 YEAST - A-AGGLUTININ	Phospholipase D. Active site motif		29331826, 35696052, 264907, 29331830,
UNCLASSIFIED gil295671 (L11275) - selected as ulant of the subunit AC40 of DNA se f and iii [Saccharomyces	_		ATTACHMENT SUBUNIT PRECURSOR			52644045, 56182435, 60432229, 264592,
gi[295671 (L11275) - selected as utant of the subunit AC40 of DNA se I and III [Saccharomyces						60433356, 60433438, 264689, 21906767.
gi[295671 (L11275) - selected as utant of the subunit AC40 of DNA se I and III [Saccharomyces	_					55811957, 35695917, 265021, 18108376,
gi[295671 (L11275) - selected as utant of the subunit AC40 of DNA se I and III [Saccharomyces						263978, 264635, 264558, 22279000
gi[295671 (L11275) - selected as utant of the subunit AC40 of DNA se I and III [Saccharomyces	1	36675305 (6047, 6048	(0)		UNCLASSIFIED	60432049, 264760, 21906769, 55811957,
gi[295671 (L11275) - selected as utant of the subunit AC40 of DNA se I and III [Saccharomyces	_					35895917, 264690, 264555, 264559
a weak suppressor of a mutant of the subunit AC40 of DNA dependant RNA polymerase I and III [Saccharomyces   cerevisiae]	_	55706629 (6049, 6050	Novel Protein sim. GBank gi[295671 (L11275) - selected as			264593, 55811576
Cefewisiae	_		a weak suppressor of a mutant of the subunit AC40 of DNA Idenmentant RNA polymerase I and III (Serrhamment			
			Cerevisiae			

22278996, 22278997, 264490, 29331825, 264111, 265007, 60170831, 265010, 87168559, 265019, 21906765, 29148627, 263967, 20281149, 20281069, 263975, 29281071, 56526486, 22279000	18108394, 222218999, 522218990, 320305200, 222718997, 222718998, 60432049, 264259, 29331822, 29331824, 66714117, 29331825, 60432289, 29331824, 66714117, 29331825, 50432289, 26331827, 29331826, 2569065, 264907, 29331830, 265009, 264900, 265007, 265008, 265009, 264910, 33657402, 264596, 21906754, 265010, 265011, 87168559, 264600, 265018, 18108351, 264682, 264683, 264769, 264764, 264288, 264685, 29148629, 3565917, 265002, 265021, 264682, 264692, 365702, 264692, 365704190, 27486261, 18108374, 55811576, 35696423, 65274791, 264639, 264558, 83373044, 18108385, 56524486, 222779002, 222279002	22278995, 22278996, 22278959, 22278995, 2264255, 29331824, 29331827, 35696052, 264908, 265008, 265009, 60170831, 21906764, 265011, 87168559, 265018, 24906768, 21906769, 29148629, 21906765, 265021, 265022, 33657109, 27486265, 264628, 264629, 18108374, 35696423, 35695855, 264638, 60170394, 22279000, 22279002, 264482, 264584	22278997, 22278999, 29331827, 264905, 264509, 264509, 264509, 264510, 264511, 264512, 8768474, 265019, 18108351, 21906768, 264534, 264690, 264693, 263869, 18108370, 264558, 22279000, 22279002, 264482		22278995, 22278997, 22278998, 22278999, 264259, 264259, 29331824, 29331824, 29331826, 29331827, 29331828, 35686052, 264908, 255018, 21906765, 21906766, 21906767, 21906768, 265924, 18108374, 264558, 56526486, 22279000, 22279002
UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	- tm7	UNCLASSIFIED
·	Contains protein domain (PF01529) - UNCLASSIFIED DHHC zinc finger domain	Contains protein domain (PF00400) - UNCLASSIFIED WD domain, G-beta repeat		Contains protein domain (PF00001) - tm7 7 transmembrane receptor (rhodopsin family)	-
3026   87643662 (6051, 6052) Novel Protein sim. GBank gij3024052 spjP97924 KARI_RAT - KALIRIN (PAM COOH-TERMINAL INTERACTOR PROTEIN 10) (P-CIP10)	JAF15184 - (AF151847) CGI-89	94231897 (6055, 6056) Novel Protein sim. GBank gi 3080521 emb CAA18650  -  (AL022599) hypothetical protein  Schizosaccharomyces  pombe	(8)	87544928 (6059, 6060) Novel Protein sim. GBank gij3757726 emb CAA18782  - (AL022727) dJ80119.1 (olfactory receptor-like protein [(hs6M1-1)) [Homo sapiens]	91677953 (6061, 6062) Novel Protein sim. GBank gil4530587]gb AAD22105.1  - (AF132000) TADA1 protein [Homo sapiens]
87643662 (6051, 6052)	94844563 (6053, 6054)				
9026	3027	3028	3029	3030	3031

22278996, 35696286, 264259, 29331824, 29331828, 264307, 29331830, 264758, 33109954, 87168474, 87168559, 265019, 264288, 21906769, 265021, 264693, 35696423, 35695855, 264636, 56182323, 83373044, 87168518				264907, 33657402, 265021	265017			264692, 264558, 18108382, 18108385, 264567		264636
synthase	struct	UNCLASSIFIED	UNCLASSIFIED	kinase		UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	glycoprotein
Contains protein domain (PF00534) - Glycosyl transferases group 1	Contains protein domain (PF00022) - struct Actin					Contains protein domain (PF00627) - UNCLASSIFIED UBA domain				
94130124 (6063, 6064) Novel Protein sim. GBank gij1019951 (U37429) - similar to Contains protein domain (PF00534) - synthase (Gycosyl transferases group 1 (Caenorhabditis elegans)	95308321 (6065, 6066) Novel Protein sim. GBank gi 5031573 ref NP_005712.1 pACTR - ARP3 (actin-related protein 3, yeast) homolog		91220692 (6069, 6070) Novel Protein sim. GBank gij3738207lemb CAA21262  - (AL031853) conserved ATP-GTP binding protein ISchizosaccharomyces pombel	91718323 (6071, 6072) Novel Protein sim. GBank gi 728837 sp P39194 ALU7_HUMAN - III! ALU SUBFAMILY SQ WARNING ENTRY III!	95307434 (6073, 6074) Novel Protein sim. GBank gij4406590 gb AAD20040  - (AF131766) Similar to Ena-VASP like protein [Homo sapiens]	85421807 (6075, 6076) Novel Protein sim. GBank gil5360093 gb AAD42865.1 AF15509 - (AF155099) NY-REN UBA domain 18 antigen [Homo sapiens]	87332257 (6077, 6078) Novel Protein sim. GBank gij4757128jemb CAB42094.1	90933517 (6079, 6080) Novel Protein sim. GBank gil4884278 emb CAB43247.1  - (AL050037) hypothetical protein (Homo sapiens)	88312357 (6081, 8082) Novel Protein sim. GBank gij3876073 emb CAB04122.1  (281505) similar to Zinc finger, C3HC4 type (RING finger); cDNA EST EMBL:D28025 comes from this gene; cDNA EST EMBL:D28024 comes from this gene; cDNA EST EMBL:D33210 comes from this gene; cDNA EST EMBL:D33411 comes from this gene; cDNA EST	85749402 (6083, 6084) Novel Protein sim. GBank gij790236 (U21156) - sarcolemmal associated protein-2 [Oryctolagus cuniculus]
94130124 (6063, 6064) h	85308321 (6065, 6066) 1	80415373 (6067, 6068)	91220692 (6069, 6070)	91718323 (6071, 6072)	95307434 (6073, 6074)	95421807 (6075, 6076) ,	87332257 (6077, 6078)	90933517 (6079, 6080)	88312357 (6081, 8082)	
3032	3033	3034	3035	3038	3037	3038	3039	3040	3041	3042

				INC. ACCICIED	25506286 R0424269 35696052 284508
3043	87773026 (6085, 6086)	3043   87773026 (6085, 6086) Novel Protein sim. GBank gij854065 emb CAA58337  -			264905, 66712502, 56182435, 55811386,
		(Aosta) Usa (numan neipesviius O)			52644296, 55811150, 35695917, 60170615,
					33657109, 18108374, 264634, 60431850
3	_	02516102 (0007 6008) Name   Decision of Charle oil (104022 (0F042276) - 0251	Contains protein domain (PF01209) - glycoprotein	hycoprotein	22278998, 22278998, 22278999, 29331824,
4400		Novel Florest Same Services (** Control of the Cont	uhiF/COO5 methyltransferase family		56182435, 264511, 265007, 60170831.
		family and serious opposed for succession			60432229, 60433356, 33109954, 18108351,
					264288, 35695917, 18108368, 18108370,
					60170394
9000		11 - State Con Alana Protein cim CBank aild Stock Blichill A 76859 11 -	Contains protein domain (PF00096) - dna ma bind	dna ma bind	284488, 264259, 35696052, 264508, 264905,
5		A DODODO NEW A 404 & profess (Homo employed	Zinc finger, C2H2 type	ı	264509, 264906, 264907, 264909, 264511,
					265006, 264591, 264593, 33109954, 264604.
	•				264764, 264683, 264288, 264766, 264768,
					21908765, 21906768, 55811957, 35695917,
			٠		27486262, 18108370, 264628, 18108374,
					35695855, 264630, 264632, 264635, 264563,
			-		264564, 264566
30.00	00000347 /6004 60001			UNCLASSIFIED.	22278999, 29331822, 29331824, 29331825,
5					29331826, 60432289, 29331827, 29331828,
					264906, 52646317, 55811957, 60432113,
					22279000, 22279002, 264482, 264564
1906	0500004 /5000 6004)			UNCLASSIFIED	264488, 22278996, 22278997, 22278999.
3			,		29331824, 29331825, 56182435, 264511,
					265008, 265009, 265011, 265017, 284766,
					21906768, 21906769, 35695917, 52844150,
					33657349, 65274791, 35695855, 264555,
					60432113, 22279000, 264566
328	_	87629419 (6095, 6096) Novel Protein sim. GBank	Contains protein domain (PF00097) - UNCLASSIFIED	UNCLASSIFIED	264102, 29148784
}	_	01458803410blAAD25962,11AF09287 - (AF092878) zinc	Zinc finger, C3HC4 type (RING		
		RING finger protein SAG [Homo saplens]	finger)		
3049		88229955 (6097, 6098) Novel Protein sim. GBank	Contains protein domain (PF01406) - UNCLASSIFIED	UNCLASSIFIED	22278997, 29331826, 264807, 264758.
		gij5454158jrefjNP_006286.1jpVARS - valyl-IRNA	IRNA synthetases class I (C)		87168559, 265018, 264448, 21906768,
		synthetase 1			265020, 33657109, 35695855, 60432113.
					22279000 20100 201000 201000 201000 201000 201000 201000 2010000 2010000 2010000 2010000 2010000 2010000 2010000 2010000
3050	97643679 (6099, 6100	3050 87643679 (6099, 6100) Novel Protein sim. GBank gil4589642 dbj BAA76843.1 -	Contains protein domain (PF00069) - kinase	kinase	264259, 29331645, 264309, 203007, 204314.
_		(AB023216) KIAA0999 protein [Homo sapiens]	Eukaryotic protein kinase domain		203019, 204209, 21300100, 203220, 204020,
					18108385, 56526486, 87168518, 22279002.
					264566
3051	1 87750599 (6101, 6102)	C			22278997, 264595, 265019, 264286, 264653,
					8/168318
3052		57108030 (6103, 6104) Novel Protein sim. GBank		dehydrogenase	264534
		gi 117528 sp[P14755 CRYL_RABIT - LAMBDA-  CRYSTALLIN			

65274572, 56181686, 22278995, 35696286, 22278998, 264259, 60432289, 265008, 265009, 60433438, 21906754, 265010, 8716859, 264603, 265018, 265019, 264763, 264288, 21906765, 21906768, 21906769, 35695917, 18108374, 35696423, 264638, 56182323, 22279000, 264563	35696286, 35696052, 29331830, 264908. 264909, 264512, 264910, 265017, 264604. 264766, 265020, 33657109, 264628. 35695855, 264636, 264564, 264566.	6424179, 65274572, 56182575, 35598260, 22278996, 22278998, 60432049, 264259, 6042426, 60432289, 60432049, 264259, 265006, 265009, 60170831, 60432229, 60431735, 6043356, 264594, 60433735, 264594, 60433736, 265011, 8168559, 21906754, 55811366, 265011, 8168559, 265019, 1108351, 264683, 264288, 264369, 264689, 21906768, 55811957, 35695917, 60170615, 33657023, 65274620, 33657109, 35695763, 60431528, 18108374, 55810764, 55811576, 35696423, 65274791, 264636, 60431850, 18108381, 56182323, 60170394, 18108385, 60432113, 264564, 264565, 264566	26448B, 224569, 18108394, 52646842, 22278997, 22278999, 22278999, 264259, 66714117, 29331826, 29331827, 35696052, 264308, 264308, 264308, 264308, 264308, 264308, 264308, 264308, 264309, 265010, 265007, 265008, 265009, 264910, 33657402, 265017, 265019, 264769, 18108351, 264762, 264763, 264764, 264769, 264769, 264689, 21906765, 21906766, 21906767, 21906768, 21906769, 26629, 18108370, 264629, 18108370, 264629, 18108381, 264631, 264634, 264634, 264634, 264634, 264634, 264634, 264634, 264634, 18108385, 264636, 264637, 264638, 18108381, 83373044, 18108385, 264567, 264567, 264566, 264568, 264567, 264568, 264637, 264638, 264634, 264637, 264638, 264634, 264635, 264567, 264568, 264568, 264567, 264568, 264568, 264567, 264568, 264568, 264567, 264568, 264568, 264567, 264568, 264568, 264567, 264568, 264567, 264568, 264568, 264567, 264568, 264568, 264567, 264568, 264568, 264567, 264568, 264568, 264567, 264568, 264568, 264567, 264568, 264568, 264567, 264568, 264568, 264568, 264567, 264568, 26458, 26458, 26458, 26458, 26458, 26458, 26458,
UNCLASSIFIED .	UNCLASSIFIED	transport	glycoprotein
			. ,.
3053   95350373 (6105, 6106)   Novel Protein sim. GBank gil]3947613 emb CAA19465.1  - (AL023828) cDNA EŜT EMBL:M89008 comes from this gene; cDNA EST yk282d3.5 comes from this gene [Caenorhabdilts elegans]	86943510 (6107, 6108) Novel Prolein sim. GBank gij1076211 pir  S50755 - hypothetical protein VSP-3 - Chlamydomonas reinhardtii	95350537 (6109, 6110) gil4680655[gblAAD27717.1 AF13294 - (AF132942) CGI-08 protein [Homo sapiens]	91861636 (6111, 6112) Novel Protein sim. GBank gij728837{sp P39194 ALU7_HUMAN - II!! ALU SUBFAMILY SQ WARNING ENTRY II!!
3053   95350373 (6105, 6106)	3054 86943510 (6107, 6108)	3055 95350537 (6109, 6110)	3056 91661636 (6111, 6112 91661636 (6111, 6112

284508 284805 264907 264908 264909	264101, 24512, 264910, 264592, 264594, 264767, 18108374, 264635, 264555, 264637, 264639, 264565, 26486	264693	22278995, 22278996, 22278997, 22278998, 22278999, 264490, 264259, 29331824, 29331825, 29331825, 265007, 60433438, 265017, 265018, 265019, 264681, 264448, 264288, 264768, 21906765, 21908766, 21906787, 21908769, 29148629, 264636, 18108385, 264563, 284567	264567	264112, 52644296, 21906768, 33657023, 263974, 18108385	264908, 265008, 18108351, 264566	18108359, 264558	52646365, 52646842, 65274572, 56182575, 56181686, 22278995, 22278996, 22278997, 22278998, 22278997, 22278998, 22278997, 264289, 264289, 64432049, 289331824, 66714117, 264208, 264630, 264308, 264686, 264686, 264687, 264687, 264689, 264689, 264689, 264689, 264689, 264689, 264699, 264891, 264999, 264899, 264639, 18108377, 55811576, 264636, 56182323, 264558, 264639, 18108385, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 22279000, 264997, 264093, 264094, 264907, 265019, 18108351, 264766, 264507, 21906765, 21906768, 264693, 2626110, 265019, 18108351, 264766, 26471069, 22279000, 222790002, 264492, 2641005, 264100, 264907, 21906768, 21906768, 264492, 2641069, 222790002, 264492, 2641069, 222790002, 264492, 264107, 264107, 2641002, 264492, 264107, 264107, 21906768, 2646902, 264492, 264107, 264107, 2641002, 264492, 264492, 264107, 2641069, 222790002, 264492, 264107, 264107, 2641002, 264492, 264107, 264107, 2641002, 264492, 2641069, 2641089, 264492, 264107, 2641089, 264
		UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	UNCLASSIFIED	transferase	UNCLASSIFIED	UNCLASSIFIED
			Coniains protein domain (PF00097) - UNCLASSIFIED Zinc finger, C3HC4 type (RING finger)					Contains protein domain (PF00403) - UNCLASSIFIED Heavy-metal-associated domain UNCLASSIFIED
	3057 95412746 (6113, 6114) Novel Protein sim. GBank gil3878119jemblCAA88860j - (Z49068) similar to GTP-binding protein; cDNA EST EMBL:M89111 comes from this gene; cDNA EST EMBL:D27709 comes from this gene; cDNA EST EMBL:D27709 comes from this gene; cDNA EST EMBL:D27708 comes from this gene; cDNA EST EMBL:D27708 comes from this gene; cDNA EST		87629425 (6117, 6118) Novel Protein sim. GBank 87629425 (6117, 6118) Novel Protein Sim. GBank 814588034[gb]AAD25962.1[AF09287 - (AF092878) zinc RING finger protein SAG [Homo sapiens]			87619465 (6123, 6124) Novel Protein sim. GBank gil4454690 gb AAD20963  - (AF070657) glutathione S-transferase subunit 13 homolog	80078023 (6125, 6128) Novel Protein sim. GBank gi 2246532 (U93872) - ORF 73, contains large complex repeat CR 73 (Kaposi's sarcoma-	91241526 (6127, 6128) Novel Protein sim. GBank gild240315[db][BAA74936.1] - (AB020720) KIAA0913 protein [Homo sapiens]  (AB020720) KIAA0913 protein [Homo sapiens]  91639201 (6129, 6130) Novel Protein sim. GBank gil5656743[gb]AAD45960.1[AC00506 - (AC005067) Supported by Human EST H08032.1 (NID:g872854), mouse EST AA870042.1 (NID:g2965487), and genscan [Homo sapiens]
	95412746 (6113, 6114)	70646238 (8445 8446)	87629425 (6117, 6118)	79346691 (6119, 6120)	1		80078023 (6125, 6126)	
	3057	900	3000	3060	3061	3062	3063	3065

18108397, 22278995, 56994075, 22278996, 264905, 66712502, 265006, 264512, 264910, 264758, 60174639, 264760, 18108351, 264764, 264683, 18108359, 264692, 18108364, 18108369, 18406370, 18108377, 18108379, 60170394, 264567	264488, 264489, 35696286, 22278996, 56994075, 264259, 29331822, 29331825, 35696052, 29331822, 29331825, 35696052, 29331822, 29331825, 2650605, 264509, 264510, 264511, 264512, 265008, 265009, 264510, 264511, 264512, 265008, 265009, 264510, 264511, 264512, 265009, 265010, 264010, 264591, 264592, 264593, 265010, 265010, 264010,	264112	22278996, 56994075, 22278998, 22278999, 264259, 264107, 264905, 29331830, 52644045, 264110, 60170831, 264592, 264594, 23657402, 21905754, 333199954, 87168474, 87168559, 265017, 264448, 264764, 264683, 264766, 52644229, 21908765, 21906768, 21906769, 264634, 284557, 60170394, 56182323, 18108385, 87168518, 22279000, 264482
UNCLASSIFIED	struct		stract
	Contains protein domain (PF00787) - struct PX domain		Contains protein domain (PF01926) - struct GTPase of unknown function
Novel Protein sim. GBank gi 4884268 emb CAB43245.1  - (AL050028) hypothetical protein [Homo sapiens]	95422551 (6133, 6134) Novel Protein sim. GBank gil4689258lgbbAAD27832.1 AF12185 - (AF121859) sorting nexin 9 [Homo sapiens]		95412753 (6137, 6138) Novel Protein sim. GBank gij3878119jembjCAA88860j - (Z4906b) similar to GTP-binding protein; cDNA EST EMBL:N89111 comes from this gene; cDNA EST EMBL:027709 comes from this gene; cDNA EST EMBL:D27708 comes from this gene; cDNA EST EMBL:D73788 comes from this gene; cDNA EST
3066   91224437 (6131, 6132) Novel Protein sim. GBank gi (AL050028) hypothetical pro	3067 95422551 (6133, 6134) N	3068 85360651 (6135, 6136)	95412753 (6137, 6138)

264488, 22278994, 22278995, 25278996, 56994075, 22278997, 22278999, 264259, 29331822, 29146498, 29146499, 66714117, 29331826, 29146498, 29146499, 66714117, 25331830, 5264445, 56182435, 264511, 265007, 264512, 264910, 60170831, 264592, 264419, 19168754, 87168474, 265019, 18108351, 264481, 264683, 264628, 21906765, 21906766, 21906766, 219067677, 21906767,	264488, 65274572, 18108398, 22278939, 35696286, 22278997, 22278998, 22278999, 264595, 264595, 264595, 264595, 264595, 264595, 264595, 29331826, 29331826, 29331826, 29331826, 294590, 264908, 29331828, 29146499, 264907, 264908, 26331820, 265007, 264512, 265008, 265009, 60170831, 60432229, 264592, 60433356, 33657402, 60433438, 33109954, 52644296, 87168474, 265010, 265017, 264681, 264296, 87168474, 265010, 265017, 264691, 265022, 264697, 265020, 266921, 265022, 264692, 33657023, 264691, 264692, 33657023, 2646937, 35694150, 264691, 264632, 264634, 135698855, 264630, 264637, 264635, 264630, 22279000, 22279000, 22799002, 264532, 264563, 264567, 264569, 264567, 264563, 264567, 264566, 264567, 264566, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264563, 264567, 264567, 264567, 264567, 264567, 264567, 264567, 264563, 264567,	56182575, 29331822, 29331824, 29331825, 29146498, 264908, 52644045, 56182435, 265009, 60433438, 55812038, 18108351, 264683, 264369, 52644150, 33657023, 264693, 33657109, 18108374, 55811576, 65274791, 284555, 56182323, 60432113, 284564
synthase	<b>5</b> 0	transcriptfactor
	Contains protein domain (PF00085) - tgf	
3070 [94319173 (6139, 6140) Novel Protein sim. GBank gij3877788 emb CAB05527 - (Z83110) cDNA EST yk472b5.3 comes from this gene; cDNA EST yk472b5.5 comes from this gene; cDNA EST yk472b5.5 comes from this gene; cDNA EST yk468c10.3 comes from this gene; cDNA EST yk468c10.5 comes from this gene; cDNA EST yk468c10.5 comes from this gene; cDNA EST yk468c10.5 comes from this gene; cDNA EST EM	94325573 (6141, 6142) Novel Protein sim. GBank gil4502425 ref NP_001709.1 pBMP6 • bone morphogenetic protein 6 precursor	95115892 (6143, 6144) Novel Protein sim. GBank gil 1263289 (U47856) - fibroin-4 [Araneus diadematus]
0 94319173 (6139, 6140) N (C)	3071 94325573 (6141, 6142) N	3072   95115892 (6143, 6144)

			**************************************	I	
3073	86147248 (6145, 6146)	3073   86147248 (6145, 6146) Novel Protein sim. GBank gij134840jspjP22528jCORB_HUMAN - CORNIFIN B (SMALL PROLINE-RICH PROTEIN IB) (SPR-IB) (14.9 KD PANCORNULIN)	-		264769
3074	88089351 (6147, 6148)	88089351 (6147, 6148) Novel Protein sim. GBank gil3419847 (AC004982) - similar to yeast hypothetical protein ybk4; similar to P38164 (PID:q588461) [Homo sapiens]		UNCLASSIFIED	264488, 265019, 264448, 264288, 21906767, 264693, 18108368, 18108370, 18108374, 264567
3075		88095752 (6148, 6150) Novel Protein sim. GBank gil4557349 refINP_000456.1 pBARD - BRCA1 associated RING domain 1	Contains protein domain (PF00023) - homeobox Ank repeat		264509, 264907, 264689, 264693, 56526486
3076	87819219 (6151, 6152)			UNCLASSIFIED	18108398, 29331822, 29331827, 60432229, 265017, 264691, 264693
3077	88734277 (6153, 6154)	88734277 (6153, 6154) Novel Protein sim. GBank gij3023956jspjQ00808jHET1_PODAN - VEGETATIBLE INCOMPATIBILITY PROTEIN HET-E-1	Contains protein domain (PF00400) - kinase WD domain, G-beta repeat	kinase	65274572, 35696052, 264511, 60170831, 87168474, 264369, 35695917, 33657182, 27486264, 33657349, 35695763, 35695855, 264639
3078		88089355 (6155, 6156) Novel Protein sim. GBank gij3900850 (AC004994) - similar to d1026456 (PID:g3043724) [Homo saplens]			22279002
3079		Novel Protein sim. GBank (Z81052) Similarity to Yea (SW:ABC1_YEAST); cDN gene; cDNA EST yk229g! [Caenorhabditis elegans]	<b>v</b>	transport	29331824, 29331826, 264758, 55811386, 265017, 55811150, 52644229, 21906768, 265020, 265021, 264693, 18108376, 264631, 52644332, 22279002
3081		B8094664 (6161, 6162) Novel Protein sim. GBank gilpzorzzi ignipatual zobos. I - gilpzorzzi ignipatual zobos. I - gilpzorzzi ignipatual zobos. I - gilpzorzzi ignipatual zobosza zobosz		UNCLASSIFIED	264486, 5264490, 6227,8999, 522,78998, 264488, 5264490, 264259, 29331827, 29331827, 29331828, 29331827, 29331828, 29331827, 29331828, 2946499, 264508, 264905, 264608, 264508, 264608, 264509, 264689, 2646445, 60174639, 264683, 264764, 264268, 264446, 264769, 264689, 264689, 264689, 264689, 264689, 264689, 264689, 264689, 264692, 33657023, 264693, 65274620, 33657109, 27488261, 35695763, 264628, 33657023, 264693, 65274620, 33657109, 27488261, 36695763, 264697, 264692, 264690, 264510, 264500, 264510, 265901, 265909, 2646908, 264909, 264510, 264511, 265099, 2646908, 2646903, 264630, 264631, 265633, 264638, 18108332, 18108332, 18108385, 264568, 264659,
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UNCLASSIFIED		UNCLASSIFIED	UNCLASSIFIED		UNCLASSIFIED	dehydrogenase
						Contains protein domain (PF00725) - dehydrogenase 1pAF07704 - (AF077049) lambda- 3-hydroxyacyl-CoA dehydrogenase
3083   88095756 (6165, 6166) Novel Protein sim. GBank gi 866241 (U29468) - C56C10.3 gene product [Caenorhabditis elegans]	87448568 (6167, 6168) Novel Protein sim. GBank gi[476774]ptr  A37475 - probable structural component p38 - borna disease virus	87795781 (6169, 6170) Novel Protein sim. GBank gi[2565057 (U80741) - CAGH44 [Homo sapiens]	87769942 (6171, 6172) Novel Protein sim. GBank gij3894189 (AC005662) - hypothetical protein [Arabidopsis thaliana]	(174)	91224441 (6175, 6176) Novel Protein sim. GBank gij335304 (AF001549) - Unknown gene groduct [Homo sapiens]	95361242 (6177, 6178) Novel Protein sim. GBank gil4689146lgblAAD27782.1IAF07704 - (AF077049) lambda- crystallin [Homo sapiens]
3083   88095756 (6165, 61	3084 87448568 (6167, 61	3085 87795781 (6169, 61	3086 87769942 (6171, 61	3087 87462988 (6173, 6174)	3088 91224441 (6175, 6	3089 95361242 (6177. 6

19522771 (6179 6180)	3000 195322371 (6179 6180) Invasi Protein sim (SBank oil 1354050 (1147024) - MEM3	UNCLASSIFIED	60424179, 52645156, 65274572, 56182575,
,	(Mis misculis)		56181686, 22278995, 35696286, 56994075.
			22278996, 22278998, 22278999, 264259.
_			29331822, 56182181, 29331824, 29331825.
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			33656970, 264906, 264908, 52644045,
		7	264828, 265006, 265007, 265008, 60170831.
	-		60432229, 60433356, 33657402, 55812038,
			264758, 21906754, 33109954, 52646317,
			55811386, 52844296, 87168474, 265011.
			87168559, 265017, 265018, 265019.
			55811150, 18108351, 264681, 264448.
		_	264288, 264369, 18108357, 264768,
			52644229, 56181562, 21906764, 21906765,
			21906766, 21906767, 21906768, 21906769,
			35695917, 265020, 265022, 60170615.
			264690, 52644150, 264691, 33657023.
			18108365, 65274620, 33657109, 18108368,
			33657182, 27486261, 27486265, 35695763,
			18108374, 18108376, 55810764, 35696423.
			55811576, 65274791, 35695855, 264557,
			56182323, 83373044, 18108387, 18108388.
			87168518, 22279000, 22279002, 264563,
		•	264482
317424 (6181, 6182)	3091   95317424 (6181, 6182) Novel Protein sim. GBank gil3873932 emb CAB01859  -	UNCLASSIFIED	35696286, 29331822, 35696052, 264508,
	(279596) Simialrity to Bovine aspartyl beta hydroxytase		264509, 264905, 264906, 264908, 264909.
	(TR:G162694); cDNA EST EMBL:D27916 comes from this		264510, 264758, 265010, 265011, 264683.
	gene; cDNA EST EMBL:027915 comes from this gene;		264685, 264768, 264768, 264769, 264693.
	cDNA EST EMBL:D64881 comes from this gene; cDNA		264628, 35696423, 35695855, 264632,
	EST EMBL: D68139 comes f		264635, 264639, 264482, 264563, 264486

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bosomalprot	UNCLASSIFIED	collagen
Contains protein G5 Ribosomal protein S5		gi400734 spiP31044 PBP_RAT - Contains protein domain (PF01161) - collagen nLAMINE-BINDING PROTEIN (23 Phosphatidylethanolamine-binding protein
2_HUMAN - 40S RIBOSOMAL	94318457 (6185, 6186) Novel Protein sim. GBank gi 5002587 emb CAB44347.1  - (Y17454) LSFR1 protein [Homo sapiens]	94316875 (6187, 6188) Novel Protein sim. GBank giļ400734 sp P31044 PBP_RAT - Contains protein domain (PF01161 PHOSPHATIDYLETHANOLAMINE-BINDING PROTEIN (23 Phosphatidylethanolamine-binding KD MORPHINE-BINDING PROTEIN) (P23K)
3092   95314592 (6183, 6184)   Novel Protein sim. GBank gli 710756[sp P15880 RS PROTEIN S2 (S4) (LLREF	3093 94318457 (6185, 6186)	3094 94316675 (6187, 6188)

5603	94848162 (6189, 6190)		Contains protein domain (PF01454) - UNCLASSIFIED MAGE family		18108397, 56182575, 22278995, 35696286. 56994075, 22278997, 22278999, 264259. 60432049, 66714117, 29331825, 60432289. 35696052, 33656970, 29146499, 264508. 264005, 284509, 29331830, 264909, 264510, 264511, 264512, 265007, 265008, 265009, 265011, 87168559, 265017, 265018, 265019, 264760, 264681, 264682, 264683, 264764, 264569, 284689, 21906765, 21906767, 265020, 265021, 265022, 264697, 265020, 265021, 265022, 264697, 265020, 265021, 265023, 264693, 264952, 18108376, 264639, 33373044, 18108385, 18108387, 264563, 264564, 264568, 284482, 264563, 264564, 264568, 284482, 264563, 264564, 264568, 284487, 18108391
3096	87756128 (6191, 6192)	87756128 (6191, 6182) Novel Protein sim. GBank gil3882221  dbj BAA34470.1  - (AB018293) KIAA0750 protein [Homo sapiens]	Contains protein domain (PF00307) - struct Calponin homology (CH) domain		22278995, 22278995, 22278997, 22278999, 29331824, 29331825, 29331826, 29331827, 33656970, 264908, 265008, 264910, 33657402, 265011, 265017, 265018, 264369, 265020, 60170615, 264691, 264692, 264693, 27486262, 18108370, 60431528, 264634, 264636, 264639, 26279000, 264566
3097	88264895 (6193, 6194 ,	88264895 (6193, 6194) Novel Protein sim. GBank gil4468288 emb CAB37981  - (AL022395) dJ273N12.1 (PUTATIVE protein based on EST matches) [Homo sapiens]	Contains protein domain (PF00646) - UNCLASSIFIED F-box domain.	UNCLASSIFIED	26488, 2931822, 29331825, 60432289, 29331826, 35696052, 29331828, 29331830, 264594, 55812038, 33109954, 33657084, 87168474, 87168559, 52644229, 21906765, 21906767, 18108378, 35696423, 52644332, 264638, 60432113, 22278002
3088	80258024 (6195, 6196	3088   80258024 (6195, 6198)			264634, 264637, 264365
3089	91243325 (6197, 6198			cyto450	26488, 35696286, 29331827, 29331829, 29331825, 29331827, 265007, 265008, 265010, 265011, 265018, 265019, 18108357, 21906766, 255020, 265022, 55811576, 56182323, 22279002, 264563
3100	87602421 (6199, 6200	3100 87602421 (6199, 6200) Novel Protein sim. GBank gij 1083764 piri B48013 - proline- rich proteoglycan 2 precursor, parolid - rat		UNCLASSIFIED	29331825, 60432289, 35696052, 264910, 60432229, 264592, 264288, 264693, 283967, 264635
3101	79602134 (6201, 6202)	n)		UNCLASSIFIED	264908, 264693, 264628, 264630, 264632

14F12853 - (AF128536) in PACSINZ [Homo saplens] 7_MOUSE - RAS-RELATED 5.1pTSSC - tumor-suppressing 6pi1938574 (U97190) - B0025.2 dits elegans] 6pi334141 emb CAA76851  - 6pi334144 emb CAA76851  - 6pi334144 emb CAA76851  - 6pi334144 emb CAA76851  - 6pi334144 emb CAA76851  - 6pi34414 emb CAA76851  - 6pi34414 emb CAA76851  - 6pi34414 emb CAA76851  - 6pi3444 emb CAA76851	1000 6000	3102   01220802 (6203 6204)   Navel Protein eim Chark	Contains profein domain (PF00018) - Struct	struct	35696286. 22278996. 22278999. 29331827,
in PACSINZ [Homo saplens]  7_MOUSE - RAS-RELATED  1.IpTSSC - tumor-suppressing  4.ip1938574 (U97190) - B0025.2  4.ip1938574 (U		=	SH3 domain		35696052, 264809, 264512, 265008,
AMOUSE - RAS-RELATED  S.IpTSSC - tumor-suppressing  gil 938574 (U97190) - B0025.2  ditis elegans)  reciption factor [Galtus gallus]  reciption factor [Galtus gallus]					60170831, 60433356, 33109954, 18108351,
7_MOUSE - RAS-RELATED 3.1 pTSSC - tumor-suppressing 3.1 pTSSC - tumor-suppressing 4 i1938574 (U97190) - B0025.2 4 i19334741 emb CAA76851  - 6 i3341441 emb CAA76851  - 6 i3341441 emb CAA76851  - 6 i3341441 emb CAA76851  - 6 i197SSC - tumor-suppressing					264684, 264689, 21906767, 60170615,
7_MOUSE - RAS-RELATED  1. pTSSC - tumor-suppressing  1. p1SSC - tumor-suppressing  1. q1938574 (U97190) - B0025.2  dits elegans   1. q13341441 emb CAA76851  -  1. q134141 emb CAA76851  -  1. q13414  -  1. q					264692, 33657023, 264638, 22279000,
### AMOUSE - RAS-RELATED  ### UNCLASSIFIED  ###################################					264482, 264564
IlpTSSC - tumor-suppressing UNCLASSIFIED (IlpTSSC - tumor-suppressing (Ilp	6205, 6206)	<u>~</u> .		UNCLASSIFIED	35695917, 264565
Ilits elegans]  Sign 3341441 [emb]CAA78851] - Secription factor [Galtus gallus]  UNCLASSIFIED  SilpTSSC - tumor-suppressing		PROTEIN RAB-17			034130 00163300 003130 030130
gil1938574 (U97190) - B0025.2 litis elegans  gil3341441 emb CAA76851 - nscription factor [Gallus gallus] 3.1 pTSSC - tumor-suppressing	(6207, 6208)	roi.		UNCLASSIFIED	. 264239, 264664, 264338, 33837 188, 264338
gi[1938574 (U97190) - B0025.2 litis elegans  litis elegans  gi[3341441 emb CAA76851  - nscription factor [Galfus gallus]	-				22278997, 264259, 29331824, 35696052,
gil1938574 (U97190) - B0025.2 litis elegans  gil3341441  emb CAA76851  - sscription factor [Galtus gallus] 3.1  pTSSC - tumor-suppressing	, , , , , , , , ,				29331828, 264508, 264509, 264905, 264908,
gi[1938574 (U97190) - B0025.2 litis elegans  gi[3341441]emb[CAA76851] - nscription factor [Galtus gallus] 3.1[pTSSC - tumor-suppressing		-			264907, 264908, 264511, 264910, 264591,
gi[1938574 (U97190) - B0025.2 litis elegans]  gi[3341441[emb]CAA76851] - scription factor [Gallus gallus]  scription factor [Callus gallus]					284594 284758 264760 264681 264762.
gi[1938574 (U97190) - B0025.2 litis elegans] gi[3341441[emb]CAA76851] - nscription factor [Galhus gallus]					264764, 264288, 264768, 264768, 264687,
gi[1938574 (U97190) - B0025.2  litis elegans]  gi[3341441]emb[CAA76851] - sscription factor [Gallus gallus]  3.1 pTSSC - tumor-suppressing					264769, 21906766, 21906768, 35695917,
gi[1938574 (U97190) - B0025.2   Itis elegans  gi[3341441 emb[CAA76851  - sscription factor [Galtus gallus]   Sscription factor [Galtus gallus]   3.1 pTSSC - turnor-suppressing					33657023, 264692, 264693, 264628, 264629,
gi[1936574 (U97190) - B0025.2 litis elegans] gi[3341441 emb CAA76851  - nscription factor [Galtus gallus] 3.1 pTSSC - turnor-suppressing					35695855, 264630, 264631, 264632, 264634,
gi[1938574 (U97190) - B0025.2   Iltis elegans]   Scription factor [Galfus gallus]   UNCLASSIFIED   UNCLASSIFIED					264635 264637 264638 264639 83373044.
gi[1938574 (U97190) - B0025.2 [itts elegans] gi[3341441[emb[CAA76851] - hscription factor [Gailus gailus] 3.1 pTSSC - turnor-suppressing	-				264404 22278002 264563 264565 264566.
gil1938574 (U97190) - B0025.2  litis elegans  gil3341441[emb CAA76851] - nscription factor [Galfus gallus] 3.1 pTSSC - turnor-suppressing					264486 264587
ilits elegans   gij3341441emb[CAA76851 - nscription factor [Gallus gallus]  UNCLASSIFIED  3.1 pTSSC - turnor-suppressing	(6211 6212)				22278996, 22278997, 22278998, 22278999,
UNCLASSIFIED UNCLASSIFIED	,				264092, 264093, 264094, 29331822, 264906,
UNCLASSIFIED UNCLASSIFIED					264907, 284908, 52644045, 56182435,
UNCLASSIFIED UNCLASSIFIED					264112, 265008, 265009, 55812038, 265017,
UNCLASSIFIED UNCLASSIFIED				g	285018, 264683, 264688, 264687, 264768,
UNCLASSIFIED UNCLASSIFIED					52644229, 21906765, 21906768, 21906769.
J UNCLASSIFIED					45811057 265020 265022 264600
J UNCLASSIFIED					53611331, 233620, 263622, 253630; 63644160, 364603, 364603, 18408370
J UNCLASSIFIED					12044130, 204032, 204033, 10100310;
UNCLASSIFIED UNCLASSIFIED					16106577, 33611376, 36162323, 16106363,
UNCLASSIFIED 99	(6213 6214)	Novel Protein eim CBank nil 3141441 lembi CAA768411			22278995 22278996 35696286 22278997.
UNCLASSIFIED UNCLASSIFIED	(	(Y17794) winged-helix transcription factor (Gallus gallus)		-	22278999, 264091, 264093, 264259,
UNCLASSIFIED 1.1pTSSC - tumor-suppressing					29331822, 29331825, 29331826, 60432289,
UNCLASSIFIED 1.1pTSSC - tumor-suppressing					29331827, 29331828, 33656970, 264105,
UNCLASSIFIED .1 pTSSC - tumor-suppressing					264512, 265009, 60433356, 60433438,
UNCLASSIFIED 3.1 pTSSC - tumor-suppressing					265011, 265017, 265018, 21906765.
UNCLASSIFIED 3.1 pTSSC - tumor-suppressing					21906768, 21906767, 21906769, 265021,
UNCLASSIFIED 3.1 pTSSC - tumor-suppressing					264691, 33657109, 27486261, 27486265,
UNCLASSIFIED 3.1 pTSSC - tumor-suppressing					18108370, 263972, 18108374, 55811576.
UNCLASSIFIED 3.1 pTSSC - tumor-suppressing					18108385, 56526486, 264482, 264487
glj5032207 ref NP_005698.1 pTSSC - tumor-suppressing	(6215, 6216)			UNCLASSIFIED	56182435, 264288, 264690, 264564
		gij5032207jrefjNP_005698.1 pTSSC - tumor-suppressing			

264490, 264908, 265007, 264910, 264593, 264683, 264684, 264687, 21906767, 21906768, 264693, 18108370, 264629, 18108374, 264632, 264638, 22279000	264488, 65274572, 22278395, 25278393, 60432049, 264259, 29331822, 29331824, 29331825, 29331825, 29331825, 29331825, 29331825, 29331825, 29331825, 29331825, 29331825, 29331825, 29331825, 295008, 265009, 6043229, 33657402, 265017, 265014, 87168559, 264600, 265017, 265018, 265019, 18108351, 264288, 265020, 60170615, 264693, 65274620, 18108370, 264639, 18108384, 22279000, 264563, 18108384, 22279000,	264488, 18108398, B6772504, A55017, 265018, 265019, 264448, 21906767, 265020, 33657023, 18108365, 18108388, 35696423, 52644332, 18108385, 18108388	ACCOUNT OF THE PERSON	18108391, 22278393, 204239, 43331824, 35696052, 264907, 264757, 60433438, 87168559, 264763, 264448, 18108354, 264288, 21906767, 21906769, 35695917, 264690, 264691, 264692, 264693, 18108365, 18108381, 18108384, 18108385, 18108388, 87168518, 22279000, 22279002	264909, 56182435, 264910, 21906754	60424179, 56182575, 264259, 29331824, 60424269, 29331828, 66712502, 264510, 265007, 60431735, 60433356, 55812038, 55811386, 265019, 264288, 264689, 21806769, 264691, 33657023, 264693, 60431528, 263974, 60431850, 56182323, 264559, 22279000, 22279002	264905, 264758, 21906764, 264690	264510, 265011, 18108351, 264288, 264689, 264689, 264689, 18108368, 18108372, 263981, 264558, 264564
struct	UNCLASSIFIED	transcriptfactor		<b>7</b> m1	- hydrolase	transcriptfactor	UNCLASSIFIED	UNCLASSIFIED
Contains protein domain (PF00096) - struct Zinc finger, C2H2 type		gij3282231 (U75454) - C2H2 type Contains protein domain (PF00096) - transcripttactor sapiens) Zinc finger, C2H2 type	Contains protein domain (F1 Cocto) Homeobox domain		Contains protein domain (PF00702) - hydrolase haloacid dehalogenase-like hydrolase			
47.1 -	95090716 (6219, 6220) Novel Protein sim. GBank gil1076211 pir IS50755 - hypothetical protein VSP-3 - Chlamydomonas reinhardiii	87754512 (6221, 6222) Novel Protein sim. GBank gij3282231 (U75454) - C2H2 type zinc finger protein [Homo sapiens]	88043639 (6223, 6224) Novel Protein sim. GBanx gij.3900846 (AC000023) - maidn   to EST AA361117 (NID:92013436) [Homo sapiens]	88207098 (6225, 6226) Novel Protein sim. GBank gi 2459910 (AF005856) -   anon2A5 [Drosophila yakuba]	79843167 (6227, 6228) Novel Protein sim. GBank gil4966270[gb]AAB52261.2] - (U97002) similar to acyl-CoA dehydrogenases and epoxide hydrolases; Pfam domain PF00441 (Acyl-CoA_dh), Score=57.4, E-value=1.7e-16, N=2; contains similarity to Pfam domain PF00702 (Hydrolase), Score=57.4, E-value=1e-13, N=1 IC	94117996 (6229, 6230) Novel Protein sim. GBank gi 5032225 ref NP_005676.1 pWBSC - Williams-Beuren syndrome chromosome region 11		
94318461 (6217, 6218)	95090716 (6219, 6220)	87754512 (6221, 6222)	88043639 (6223, 6224)	88207098 (6225, 6226)	79843167 (6227, 6228)		79642855 (6231, 6232)	87771288 (6233, 6234)
3109	3110		3112	3113	3114	3115	3116	3117

118   94655846 (6235, 6239)   Novel Protein sim. GBank gij280565]pmb[CAB01444.11 -   Contains protein domain (PF00008) - (gr. (12010) period to using demendent similar to enrichtly period (230, 6239)	52645156, 52646842, 65274572, 56182575, 22278995, 56994075, 22278996, 35696286, 22278995, 22278996, 35696286, 22278997, 22278999, 22278999, 2569286, 22278997, 22278999, 2569286, 29331827, 256908, 256331826, 29331827, 25690652, 29331828, 2644045, 56182435, 264510, 264511, 265007, 265008, 265009, 264757, 52646317, 21906754, 33657084, 5264428, 8716859, 265017, 265018, 26469, 264005, 265019, 264762, 264448, 265018, 26469, 264005, 264005, 264007, 265019, 264762, 264691, 33657023, 264693, 33657109, 33657349, 264628, 18108376, 60431528, 81088374, 87188518, 22279000, 22279002, 264486	265006, 264288	264488, 264509, 264510, 264511, 264512, 264288, 264486	5264507, 52645156, 52646365, 52646842, 22278994, 56994075, 22278996, 22278999, 264259, 29331824, 29331827, 35696052, 52644045, 265008, 528461317, 87168474, 87168559, 21906765, 52844150, 33857023, 18108374, 264637	264638	18108392, 29331822, 29331824, 29331825, 264905, 265007, 55812038, 265019, 18108351, 264682, 264288, 264766, 21906764, 21906765, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906768, 21906769, 25811957, 18108381, 18108384, 22279000, 22279002, 284482	264905	56181686, 264259, 66714117, 60432289, 29331826, 29331827, 264907, 264908, 264828, 265009, 60433356, 33657402, 60433438, 264758, 18108351, 264288, 29148827, 29148629, 33657023, 33657109, 18108382, 56526488
pij3880563jembjCAB0144.1j - (Senefinder, similar to DNA EST yk353d10.5 comes from slegans]  1/AF14315 - (AF143152) putative uplex I subunit jCeenorhabditis gij2439517 (AC002563) - putative gij2439517 (AC002563) - putative gij2439517 (AC002663) - putative gij2439517 (AC001714) in dehydrogenase/reductase naj	jō		UNCLASSIFIED		UNCLASSIFIED	kinase	UNCLASSIFIED	dehydrogenase
665848 (6235, 6238) Novel Protein sim. GBank gij3890563jemb CAB01444.1 - (Z78018) predicted using Genefinder; similar to serine/threonine kinase; cDNA EST yk353d10.5 comes from this gene [Caenorhabditis elegans]  344040 (6239, 6240) Novel Protein sim. GBank gij383016. (AF14315) putative NADH oxtdoreductase complex I subunit [Caenorhabditis elegans]  110735 (6241, 6243, 6244) Novel Protein sim. GBank gij2439517 (AC002563) - putative RHO/RAC effector protein; 95% similarity to P49205 (PID:g1345860) [Homo sapiens]  788899 (6245, 6246) Novel Protein sim. GBank gij2439517 (AC002563) - putative RHO/RAC effector protein; 95% similarity to P49205 (PID:g1345860) [Homo sapiens]  1216607 (6249, 6250) Novel Protein sim. GBank gij2439517 (AC001714) oxidoreductase family [Thermotoga martitima]	Contains protein domain (PF00008) -	Contains protein domain (PF00328) - Histidine acid phosphatase				Contains protein domain (PF00780) - CNH domain		Contains protein domain (PF00106) - short chain dehydrogenase
665848 (6235, 6236) 728796 (6237, 6238) 344040 (6239, 6240) 110735 (6241, 6242) 683003 (6245, 6246) 786899 (6247, 6248) 7116607 (6249, 6250)	gij3880563jembjCAB01444.1 - Senefinder, similar to NA EST yk353d10.5 comes from elegans		Novel Protein slm. GBank gi 5019819 gb AAD37863.1 AF14315 - (AF143152) pulative NADH oxidoreductase complex I subunit  Caenorhabditis elegans	Novel Protein sim. GBank gi 4501877 ref NP_001088.1 pACR  - acrosin		Novel Protein sim. GBank gi[2439517 (AC002563) - putative RHO/RAC effector protein; 95% similarily to P49205 (PID:g1345860) [Homo sapiens]		Novel Prolein sim. GBank gil4980826[gb]AAD35412.1 AE00171 - (AE001714) oxidoreductase, short chain dehydrogenase/reductase family [Thermotoga maritima]
3118 94 3120 87 3122 11 3123 88 3123 88 3124 88	94665848 (6235, 6236)		87344040 (6239, 6240)	94110735 (6241, 6242)	11814528 (6243, 6244)	88083003 (6245, 6246)	87786899 (6247, 6248)	91216607 (6249, 6250)

	3126   95337205 (6251, 6252)			UNCLASSIFIED	22278989, 264490, 264259, 60432049, 29331822, 60432289, 29146498, 52644045,
	_		,	-	56182435, 265009, 60433438, 265010,
					8/168559, 265017, 265016, 55811150, 264763, 264683, 264369, 264685, 29148629,
					33657023, 264693, 33657109, 18108374,
_			,		55811576, 18108385, 60432113, 22279002
-	91639233 (6253, 6254)	91639233 (6253, 6254) Novel Protein sim. GBank gi[2828280 emb[CAA16694.1] -			35696286, 22278996, 22278999, 29331828,
_	,				264908, 60433438, 87168559, 264604,
_					21906765, 21906769, 33657023, 33657349,
					264629, 18108374, 18108377, 22279000.
					22279002
_	87674330 (6255, 6256)	3128 187674330 (6255 6256) Novel Protein sim. GBank gil3885828 (AF090133) - lin-7-A	Contains protein domain (PF00595) - misc channel	misc channel	22278996, 264259, 52644045, 265008,
_			PDZ domain (Also known as DHR or	1	21906754, 265017, 265018, 21906768,
			GLGF).		18108376, 18108387, 22279000, 22279002
	87755412 (6257, 6258)	3129   87755412 (6257, 6258) Novel Protein sim. GBank gil3135273 (AC003058) -	Contains protein domain (PF00400) - kinase	kinase	56182575, 264259, 29331825, 29331828,
			WD domain, G-beta repeat		52644045, 56182435, 60433356, 264600,
					264682, 264763, 264764, 264369, 264288,
					264686, 55811957, 264692, 33657023,
					33657109, 60432113, 264564, 264566
	14993960 (6259, 6260)	14993960 (6259, 6260) Novel Protein sim. GBank gij3329465 (AF064553) - NSD1			264636
		protein [Mus musculus]			
	95351469 (6261, 6262)	95351469 (6261, 6262) Novel Protein sim. GBank gi 1846277 (U86136) -	Contains protein domain (PF00400) - UNCLASSIFIED	UNCLASSIFIED	56182575, 264259, 29331824, 264907,
		telomerase-associated protein TP-1 [Homo sapiens]	WD domain, G-beta repeat		56182435, 264594, 60433438, 55812038,
					33109954, 21906754, 33657084, 87168474,
					264448, 264768, 21906769, 55811957,
					265020, 265021, 265022, 60170615,
					33657023, 33657109, 33657182, 27486261,
_	٠				33657349, 65274791, 60170394, 56182323.
					83373044, 87168518, 264564

52644507, 52646842, 52646365, 65274572, 56182575, 22278994, 22278995, 35696286, 56994075, 22278994, 22278995, 25278998, 22278998, 22278998, 22278998, 22278998, 22278998, 22278998, 22278998, 22278999, 22278998, 22278999, 29331824, 29331825, 60432289, 29331826, 29331826, 25618289, 26331826, 25618052, 264310, 6043229, 60433356, 33657402, 55812038, 52646317, 21906754, 52644296, 85658542, 87168559, 265017, 265018, 256019, 264488, 264288, 2645129, 265027, 250016765, 21906766, 21906766, 21906766, 21906766, 21906769, 3565021, 60170615, 3265109, 33657192, 27488261, 27488262, 33657349, 35695763, 18108336, 18108337, 56526486, 87168518, 60432113, 22279002	22278994, 22278998, 264909, 265006, 265007, 87168559, 264760, 21906767, 18108374, 22279000, 22279002, 264563	284595, 264369, 264669, 264626, 264500	22278996, 264095, 29331826, 33557402, 18108348, 253974	22278999, 264259, 264626, 265000, 265000, 60433438, 265019, 264764, 264288, 264769, 264689, 265020, 27486262, 263972, 65274791, 264557, 264558	22278995, 22278996, 22278994, 22278996, 22278996, 22278996, 22278996, 22278999, 264259, 29331822, 29331825, 29331825, 29331826, 29331825, 29331827, 29331827, 264510, 265008, 21906754, 87168474, 265019, 87188559, 265017, 265008, 21906765, 21906765, 21906765, 21906766, 21906769, 55811957, 35695917, 265020, 265021, 52644150, 18108370, 18108374, 22279000, 22279002, 264486, 264486
. ubiquitin	polymerase		struct	transport	UNCLASSIFIED
Contains protein domain (PF00789) - Ubiquitin UBX domain				Contains protein domain (PF00153) - transport Mitochondrial carrier proteins	
3132 95415459 (6263, 6264) Novel Protein sim. GBank gll4680647lgbbAD27713.1lAF13293 - (AF132938) CGI-03 1 protein [Homo sapiens]	3133 87379414 (6265, 6266) Novel Protein sim. GBank gil4507613 ref NP_003738.1 pTNKS - TANKYRASE	94649816 (6267, 6268) Novel Protein sim. GBank gi[1729827 sp P54633 TALA_DICDI - FILOPODIN (TALIN HOMOLOG)	n sim. GBank nding protein	94845839 (6271, 6272) Novel Protein sim. GBank gil627101  piri 544092 - probable carrier protein c2 - Caenorhabdilis elegans	88257947 (6273, 6274) Novel Protein sim. GBank gij3342730 (AC005331) - R31341_1 [Homo sapiens]
95415459 (6263, 6264)   h	87379414 (6265, 6266)	94649816 (6267, 6268)	86389356 (6269, 6270)	94845839 (6271, 6272)	88257947 (6273, 6274)
3132	3133	3134	3135	3136	3137

264569, 264488, 264907, 264511, 264593, 33109954, 87168559, 264681, 264684, 264685, 264687, 264768, 264688, 264689, 264691, 264692, 264693, 33657109, 264631, 264634, 264635, 264638, 1810838, 60170394, 83373044, 18108385, 18108388, 60432113, 22279000, 22279002	22278997, 22278986, 22278999, 264905, 265018, 265019, 21906765, 265020, 264636, 264557	22278995, 56994075, 35696286, 264908, 264909, 264909, 60433356, 21906754, 52644296, 87168474, 87168559, 264683, 264288, 264685, 264693, 27486262, 35695655, 264630, 264555, 264556	56182575, 35696286, 29331828, 264909, 265009, 265018, 18108351, 264369, 21906766, 29148627, 265020, 264628, 264629, 264631, 18108385	52645156, 52646365, 22278995, 35696286, 22278998, 22278999, 60432049, 264259, 29331822, 29331822, 29331822, 29331822, 29331822, 29331822, 29331822, 29331822, 29331822, 20331822, 2033229, 33657402, 264595, 60433438, 264766, 264687, 52644229, 21906765, 21906767, 21906768, 60170615, 52644150, 65274620, 33657499, 35695763, 18108370, 18108376, 65274791, 35695855, 264631, 264557, 87168518, 60433113, 222789000	264488, 56182575, 224218399, 22476939, 22278999, 29331822, 29331824, 60432289, 35596052, 29331822, 295608, 264905, 264907, 264908, 264909, 264907, 264908, 264508, 264910, 6043229, 33657402, 60433356, 60433438, 264448, 264764, 264684, 264760, 264763, 26468, 264769, 264689, 264760, 264689, 264760, 264689, 264280, 264680, 33657023, 264693, 263967, 33657109, 264628, 264639, 18108374, 263976, 55611576, 35695855, 264636, 264637, 264607, 264588, 87168518, 60432113, 264638, 264563, 264566, 264488
	UNCLASSIFIED	stract	UNCLASSIFIED	cytochrome	ONGLASSIFIED
					,
- l070] - aliana]	32C - Hyp-	3] - varticle	39) - putative	15,0561_PIG	major]
	Novel Protein sim. GBank gi 228938 prf  1814452C - Hyp- rich glycoprotein [Zea diploperennis]		Novel Protein sim. GBank gij3213227 (AF035209) - putative v-SNARE VII1a [Mus musculus]	tovel Protein sim. GBank gilz498197 sp Q95245 C561_PIG	Novel Protein sim. GBank gi \$420387 emb CAB46679.1  (AJ243459) proteophosphoglycan [Lelshmania major]
3138   94130186 (6275, 6276)   Novel Protein sim. GBank (AC006838) hypothetical p	87325503 (6277, 6278) Novel Pro	91222692 (6279, 6280) Novel Protein sim. GBank (X53744) 68kDA subunit o [Canis famillaris]	87323564 (6281, 6282) Novel Protein sim. GBank v-SNARE VII1a (Mus mus	95419028 (6283, 6284) Novel Protein sim. GBank - CYTOCHROME B561 (C	95351475 (6285, 6286) Novel Protein sim. GBanl (AJ243459) proteophosp
3138 94-	3139 87	3140	3141 87		243 643

	95336329 (6287, 6288)	3144   95336329 (6287, 6288)   Novel Protein sim. GBank gi 4884468 emb CAB43322.1  - (AL050225) hypothetical protein [Homo sapiens]			264488, 18108398, 22278996, 35696286, 22278997, 22278999, 29331826, 29331827, 35696052, 29331828, 264106, 265006,
					265007, 265009, 33657402, 85658542, 265011, 18108351, 284448, 264369, 21906765, 21906767, 265020, 265021, 52644150, 27486261, 18108370,
				,	18108374, 35696423, 56182323, 83373044, 22279000, 22279002, 264567
3145 86	611657 (6289, 6290)	86611657 (6289, 6290) Novel Protein sim. GBank gij3879709jemb CAB03330j - (281118) Similarity to Human endosomal protein P162		UNCLASSIFIED	18108397, 29331824, 29146499, 20281100, 265006, 55812038, 265010, 21906768,
		(TR:Q15075); cDNA EST EMBL.Z14487 comes from this gene; cDNA EST EMBL.Z14556 comes from this gene; cDNA EST EMBL:D27011 comes from this gene; cDNA EST EMBL:D27015 comes from t			29148621, 21905703, 29146764, 204032, 33657023, 33657109, 35695763, 263981, 56182323, 87168518
66	756314 (6291, 6292)	87756314 (6291, 6292) Novel Protein sim. GBank gij2135746 pir  S69890 - mitogen   Contains protein domain (PF00169) - struct   PH domain   Inducible gene mig-2 - human   PH domain   PH d	Contains protein domain (PF00169) - Is PH domain		264259, 29331626, 29331630. 264510, 264511, 265007, 265009, 264600. 265017, 18108351, 264448, 264369. 21906766, 265021, 264692, 33657109, 18108374, 35696423, 35695855, 60432113,
8	1848512 (6283, 6294 <u>)</u>	94848512 (6293, 6294) Novel Protein sim. GBank gij3874279jembjCAB07315.1j - (292825) predicted using Genefinder; cDNA EST yk315e12.3 comes from this gene; cDNA EST yk315e12.5 comes from this gene [Caenorhabditis elegans]	Contains protein domain (PF00702) - UNCLASSIFIED haloacid dehalogenase-like hydrolase	UNCLASSIFIED	56181686, 35696286, 60432049, 264259, 56182181, 29331825, 60432289, 35696052, 56182435, 265008, 264910, 60431735, 60433356, 60433438, 265010, 264448, 264288, 265022, 33657023, 33657109, 60431528, 65274791, 264631, 56182323,
8	362169 (6295, 6295 <sub>,</sub>	3148   95362169 (6295, 6296) Novel Protein sim. GBank gij5225322[gb]ΑΑΔ40851.1 AF08310 - (AF083108) sirtuin type 3 [Homo sapiens]		UNCLASSIFIED	35696286, 35696052, 264511, 85658542. 87168474, 284764, 35696423, 264555. 264556, 264557, 264558, 83373044. 56526488, 60432113
3149 9	5308548 (6297, 6298	95308548 (6297, 6298) Novel Protein sim. GBank gil4200446 (AF102777) - FYVE finger-containing phosphoinositide kinase [Mus musculus]	Contains protein domain (PF01363) - eph FYVE zinc finger	eph	29331822, 35696052, 264109, 29148629, 18108381
3150 87	7655472 (6299, 6300	87655472 (6299, 6300) Novel Protein sim. GBank gij3378454 emb CAA76893  - (Y17850) ganglioside-induced differentiation associated protein 1 [Mus musculus]	Contains protein domain (PF00043) - transferase Glutathione S-transferases.	transferase	264259, 29331822, 29331824, 29331825, 29331827, 52646317, 264688, 35695855, 56182323, 264639
18	772355 (6301, 6302	87772355 (6301, 6302) Novel Prolein sim. GBank gi[172591 (M63577) - SFP1 [Saccharomyces cerevisiae]	Contains protein domain (PF00096) - oncogene Zinc finger, C2H2 type	оисодене	29331822, 265008
3152 8	85698108 (6303, 6304)			UNCLASSIFIED	21906754, 87168559, 264605, 21906768, 52644150, 27486264, 35696423, 22279000

			(900000) -:		264488 52646365 35696286, 22278996,
153	95317299 (6305, 6306)	3153 95317299 (6305, 6306) Novel Protein sim. GBank	WD domain. G-beta repeat		22278997, 22278999, 60432049, 264259,
		032/02.1		2	29331826, 60432289, 33656970, 264508,
		3 [Mus musculus]		.5	264908, 33657402, 264595, 60433438,
		•		80	87168474, 87168559, 264601, 265019,
	-			2	264448, 264682, 264764, 264288, 264369.
_				2	264768, 21906765, 21906766, 21906767,
				8	21906768, 21906769, 29148784, 265021.
				7	265022, 60170615, 52644150, 264690,
				7	284691, 33657023, 65274620, 33657109.
					18108370, 35695855, 264638, 60170394.
					87168518, 60432113, 22279000, 22279002
_		See	A	ATPase_associated 2	22278998, 264259, 29331824, 66712502,
3154	87718573 (6307, 6308)	87718573 (6307, 6308) Novel Protein Sim. Openia			265008, 265010, 265017, 18108354, 264691.
		3		.,	33657023, 264693, 20281149, 18108374
		protein (Homo Saprens)	D	UNCLASSIFIED	29331828, 264509, 264905, 264908, 264510.
3155		87762394 (6309, 6310) Nover Protein Sim. Codula			264511, 264512, 33657402, 264681, 264683.
		gilt 2003 i japin 39 i 34 photo _ i i i i i i i i i i i i i i i i i i		-	33657023, 18108370, 264634, 264639.
		SQ WARNING ENTRY IIII			18108385, 264563, 264486
			Cantaine protein domain (PE00652) - Iransferase		56182575, 22278996, 22278997, 22278998,
3156	87737449 (6311, 6312)		Similarity to lectin domain of right		22278999, 60432049, 264259, 29331822,
		g  5630078  gp  AAD45821.1  AC00001 - (AC0000117) 14	Lots chain 3 cooles		29331824, 66714117, 29331825, 29331826.
		acetylgalactosaminyltransferase; similar to Q10473	beta-chain, a copies.		29331827, 35696052, 52644045, 265007,
_		((PID:g1709559) [Homo saplens]			256000 E0170831 E0432229 E0433356
					200003, 00110031, 00401112, 00:0010;
					219U6/34, 351U9934, 671U6414, 203010.
					265017, 265018, 265019, 16106351, 204449.
					264288, 264689, 21906766, 21906768,
					21906769, 35695917, 265020, 265022,
				•	264692, 18108370, 35696423, 56182323,
					22279002
	-1				18108396, 264259, 29331826, 35696052,
7015	88259577 (6313, 6314)				29146498, 87168559, 265017, 264448,
					264288, 264691, 18108366, 52645129,
	,				35696423, 52644332
94.60	-	20024448 /6245   6248   Novel Protein elm GBank	Contains protein domain (PF00023) - kinase	kinase	264488, 263974
2	_	gil5306084[gb]AAD41895.1[AF15677 - (AF156778) ASB-3			
		protein [Homo sapiens]			56400575 20078000 20331824 264106
3159	•	94124114 (6317, 6318) Novel Protein sim. GBank gij5531272 emb CAB50897.11 -		UNCLASSIFIED	00104270, 22270393, 23031024, 207100,
	_	(AJ243800) WSC4 homologue [Kluyveromyces lactis]			504435356, 204736, 2030111, 01 100336.
					264448, 18108354, 264768, Z19U6706.
_					265020, 264691, 264692, 3365/109.
					18108374, 35696423, 264555, 60170394.
					22279000
2460	_	90221068 (5319 6320) Novel Protein sim GBank 0 3930525 (AF064447) - sex-	Contains protein domain (PF00023) - struct	struct	18108351, 264555, 264556, 264557, 264558.
200			Ank repeat		264559

264259, 29331822, 60432289, 29331828,	52644045, 285017, 265018, 264448, 264286	21906764, 21906767, 265020, 18108374,	264636, 264566	
88074111 (6321, 6322)				
	88074111 (6321, 6322)	88074111 (6321, 6322) 264259, 29331822, 60432289, 29331828.	88074111 (6321, 6322) 2264259, 29331822, 60432289, 29331828. 52644045, 285017, 265018, 264288. 21906764, 21906767, 265020, 18108374.	264259, 29331822, 60432289, 29331828. 52644045, 285017, 265018, 26448, 264288. 21906764, 21906767, 265020, 18108374. 264636, 264566

Table 2

Tissue iD	Tissue Name	Tissue Information	Disease Association
20281069	192xN	Protein-protein Interactions	Any
20281071	192xN	Protein-protein Interactions	Any
20281149	192xN	Protein-protein Interactions	Any
20281152	192xN	Protein-protein Interactions	Any
264111	276xN	Protein-protein Interactions	Any
264112	276xN	Protein-protein Interactions	Any
263966	384xN	Protein-protein Interactions	Any
263967	384xN	Protein-protein Interactions	Any
	552xN	Protein-protein Interactions	Any
18108379	5PH 52.1 (Adrenal Gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia,
	5PH 52.2 (Fetal Lung)	Fetal Lung	Cystic Fibrosis, infection, lung cancer
18108383	5PH 52.3 (B's Lyphoma-Raji)	Lymphoma derived from B cells	Blood cancers, hematopoeisis, leukemia
18108368	5PH 52.5 (Salivary Gland)	Salivary Gland	Dry mouth, infection
18108384	5PH 52.6 (Brain- Thalmus)	Thalamus	Brain cancer, head injury, obesity, neurological disorders, neuropsychiatric disorders
18108394	5PH 53.1 (Adrenal Gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia,
18108355	5PH 53.2 (Fetal Lung)	Fetal Lung	Cystic Fibrosis, infection, lung cancer
18108359	5PH 53.3 (B's Lyphoma- Raji)	Lymphoma derived from B cells	Blood cancers, hematopoeisis, leukemia
18108361	5PH 53.4 (Mammary Gland)	Mammary Gland	Lactation disorders, breast cancer
18108362	5PH 53.5 (Salivary Gland)	Salivary Gland	Dry mouth, infection
18108366	5PH 53.6 (Brain- Thalmus)	Thalamus	Brain cancer, head injury, obesity, neurological disorders, neuropsychiatric disorders
18108354	5PH 54.1 (Adrenal Gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia,
18108392	5PH 54.2 (Fetal Lung)	Fetal Lung	Cystic Fibrosis, infection, lung cancer
18108348	5PH 54.3 (B's Lyphoma- Raji)	Lymphoma derived from B cells	Blood cancers, hematopoeisis, leukemia
18108382	5PH 54.4 (Mammary Gland)	Mammary Gland	Lactation disorders, breast cancer
18108395	SPH 54.5 (Salivary Gland)	Salivary Gland	Dry mouth, infection
18108365	5PH 54.6 (Brain- Thalmus)	Thalamus	Brain cancer, head injury, obesity, neurological disorders,
			neuropsychiatric disorders
18108397	SPH 55.1 (Adrenal Gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy, Congenital Adrenal Hyperplasia,
18108398	5PH 55.2 (Fetal Lung)	Fetal Lung	Cystic Fibrosis, infection, lung cancer
18108364		Lymphoma derived from B cells	
18108388	SPH 55.4 (Mammary Gland)	Mammary Gland	Lactation disorders, breast cancer
18108358	5PH 55.5 (Salivary Gland)	Salivary Gland	Dry mouth, infection
20281099	5PH 56.2 (MG63)		
20281100	5PH 56.3 (UISMC)		
264404	5PH.1 (Brain)	Whole Brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's disease, Stroke, Tuberous sclerosis, hypercalceimia, Parkinson's disease, Huntington's disease, Cerebral palsy, Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxiatelangiectasia, Leukodystrophies, Behavioral disorders,
<b>.</b> .			Addiction, Anxiety, Pain, Neuroprotection

264510	5PH.10 (Pancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
264511	5PH.11 (Placenta)	Placenta	Infertility, birth defects
264512	5PH.12 (Thyroid)	Thyroid	Hyperparathyroidism, Hypoparathyroidism
264555	5PH.13 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic thrombocytopenic purpura, autoimmume disease, allergies,
~			immunodeficiencies, transplantation, Graft vesus host,
264556	5PH.14 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies, immunodeficiencies, transplantation, Graft vesus host,
264557	SPH.15 (Bone Marrow)	Воле Магтом	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies, immunodeficiencies, transplantation, Graft vesus host,
264558	5PH.16 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies, immunodeficiencies, transplantation, Graft vesus host,
264559	SPH.17 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies, immunodeficiencies, transplantation, Graft vesus host,
264569	5PH.19 (One Fetal tissue and two cell lines)	Mixed	
264687	5PH.19.1 (fetal thymus - CRL7046)	Fetal Thymus	Hemophilia, hypercoagulation, Idiopathic thrombocytopenic purpura, immunodeficiencies
264688	5PH.19.2 (hematopoetic stem	Hematopoeitic stem cells	Leukemia, osteoporosis, post-chemotherapeutic stem cell
207000	cells - CRL2043)		repopulation
264689	5PH.19.3 (osteogenic sarcoma cell lines - HTB96)	Osteogenic Sarcoma	Sarcomas, osteoporosis, osteopetrosis
264690	5PH.19.4 (Fetal Liver)	Fetal liver	Von Hippel-Lindau (VHL) syndrome, Cirrhosis, Transplantation
264691	SPH.19.5 (Heart)	Heart	Cardiomyopathy, Atherosclerosis, Hypertension, Congenital heart defects, Aortic stenosis, Atrial septal
			defect (ASD), Atrioventricular (A-V) canal defect, Ductus
			arteriosus, Pulmonary stenosis, Subaortic stenosis,
	*		Ventricular septal defect (VSD), valve diseases, Tuberous sclerosis, Scleroderma, Obesity, Transplantation
264692	5PH.19.6 (Spleen)	Spleen	Hemophilia, Hypercoagulation, Idiopathic
			thrombocytopenic purpura, Immunodeficiencies, Graft vesus host
24402	5011 10 7 (Discipano)	Dia ilea	Von Hippel-Lindau (VHL) syndrome , Alzheimer's
264693	5PH.19.7 (Pituitary)	Pituitary	disease, Stroke, Tuberous sclerosis, hypercalceimia, Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection
264482	5PH.2 (Brain)	Brain	Von Hippel-Lindau (VHL) syndrome , Alzheimer's
	<b>\</b>		disease, Stroke, Tuberous sclerosis, hypercalceimia, Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
	1		sclerosis, Ataxia- telangiectasia, Leukodystrophies, Behavioral disorders,
		·	Addiction, Anxiety, Pain, Neuroprotection

264600	5PH.21 (Fetal Brain)	Fetal brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's disease, Stroke, Tuberous sclerosis, hypercalceimia, Parkinson's disease, Huntington's disease, Cerebral palsy, Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxiatelangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection
264601	5PH.22 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic thrombocytopenic purpura, autoimmume disease, allergies, immunodeficiencies, transplantation, Graft vesus host,
264602	5PH.23 (Thyroid)	Thyroid	Hyperthyroidism and Hypothyroidism
264603	5PH.24 (Pancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
264604	5PH.25 (Lymph Node)	Lymph Node	Lymphedema, Allergies
264605	5PH.26 (Placenta)	Placenta	Infertility, birth defects
264634	5PH.28 (Heart)	Heart	Cardiomyopathy, Atherosclerosis, Hypertension, Congenital heart defects, Aortic stenosis, Atrial septal defect (ASD), Atrioventricular (A-V) canal defect, Ductus arteriosus, Pulmonary stenosis, Subaortic stenosis, Ventricular septal defect (VSD), valve diseases, Tuberous sclerosis, Scleroderma, Obesity, Transplantation
264635	5PH.29 (Fetal Kidney)	Fetal Kidney	Diabetes, Autoimmune disease, Renal artery stenosis, Interstitial nephritis, Glomerulonephritis, Polycystic kidney disease, Systemic lupus erythematosus, Renal tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch-Nyhan syndrome
264483	5PH.3 (Bone Marrow)	Bone marrow	Hemophilia, hypercoagulation, Idiopathic thrombocytopenic purpura, autoimmume disease, allergies, immunodeficiencies, transplantation, Graft vesus host,
264636	5PH.30 (Lymph Node)	Lymph Node	Lymphedema , Allergies
264637	5PH.31 (P)ancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
264638	5PH.32 (Thyroid)	Thyroid	Hyperthyroidism and Hypothyroidism
264639	SPH.33 (Fetal Brain)	Fetal brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's disease, Stroke, Tuberous sclerosis, hypercalceimia, Parkinson's disease, Huntington's disease, Cerebral palsy, Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxiatelangiectasia, Leukodystrophics, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection
264484	SPH.4 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic thrombocytopenic purpura, autoimmume disease, allergies, immunodeficiencies, transplantation, Graft vesus host,
264758	5PH.44.1 (Kidney)	Kidney	Diabetes, Autoimmune disease, Renal artery stenosis, Interstitial nephritis, Glomerulonephritis, Polycystic kidney disease, Systemic lupus erythematosus, Renal tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch-Nyhan syndrome
264760	5PH.44.2 (Fetal Liver)	Fetal Liver	Von Hippel-Lindau (VHL) syndrome, Cirrhosis, Transplantation

264762	5PH.44.3 (Heart)	Heart	Cardiomyopathy, Atherosclerosis, Hypertension, Congenital heart defects, Aortic stenosis, Atrial septal defect (ASD), Atrioventricular (A-V) canal defect, Ductus arteriosus, Pulmonary stenosis, Subaortic stenosis, Ventricular septal defect (VSD), valve diseases, Tuberous sclerosis, Scleroderma, Obesity, Transplantation
264764	5PH.44.4 (Prostate)	Prostate	Prostate Cancer
264766	5PH.44.5 (Spleen)	Spleen	Hemophilia, Hypercoagulation, Idiopathic thrombocytopenic purpura, Immunodeficiencies, Graft vesus host
264768	5PH.44.6 (pituitary)	Pituitary	Von Hippel-Lindau (VHL) syndrome, Alzheimer's disease, Stroke, Tuberous sclerosis, hypercalceimia, Parkinson's disease, Huntington's disease, Cerebral palsy, Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxiatelangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection
264769	5PH.44.7 (Uterus)	Uterus	Infertility, birth defects
264905	5PH.48.1 (Burkitt's Lymphoma- Raji)	Burkitt's Lymphoma	Lymphoma, blood cancers
264906	5PH.48.2 (Thalamus- Brain)	Thalamus	Von Hippel-Lindau (VHL) syndrome, Alzheimer's disease, Stroke, Tuberous sclerosis, hypercalceimia, Parkinson's disease, Huntington's disease, Cerebral palsy, Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxiatelangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection
264907	5PH.48.3 (Adrenal Gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia,
264908	5PH.48.4 (Fetal Lung)	Fetal Lung	Cystic Fibrosis, infection, lung cancer
264909	5PH.48.5 (Salivary Gland)	Salivary Gland	Dry mouth, infection
264910	5PH.48.6 (Mammary Gland)	Mammary Gland	Lactation disorders, breast cancer
265006	5PH.50.1 (B's lymphoma)	Burkitt's Lymphoma	Lymphoma, blood cancers
265007	SPH.50.2 (thalamus)	Thalamus	Von Hippel-Lindau (VHL) syndrome, Alzheimer's disease, Stroke, Tuberous sclerosis, hypercalceimia, Parkinson's disease, Huntington's disease, Cerebral palsy, Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxiatelangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection
265008	SPH.50.3 (adrenal gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy, Congenital Adrenal Hyperplasia,
265009	5PH.50.4 (fetal lung)	Fetal Lung	Cystic Fibrosis, infection, lung cancer
265010	5PH.50.5 (salivary gland)	Salivary Gland	Dry mouth, infection
265011	5PH.50.6 (mammary gland)	Mammary Gland	Lactation disorders, breast cancer
18108385	5PH.51.1 (MCF-7)	Breast Cancer	Breast Cancer
18108370	5PH.51.2 (CCRF-CEM)	Cancer Cell line	Cancer
18108374	5PH.51.3 (K-562)	Cancer Cell line	Cancer
18108351	SPH.51.4 (OVCAR-3)	Ovarian cancer	Ovarian cancer Cancer
18108372	5PH.51.5 (HL-60)	Cancer Cell line	
264486	5PH.6 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic thrombocytopenic purpura, autoimmume disease, allergies, immunodeficiencies, transplantation, Graft vesus host,

264508	5PH.8 (Fetal Brain)		Von Hippel-Lindau (VHL) syndrome, Alzheimer's
	,		disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264509	5PH.9 (Lymph Node)	Lymph Node	Lymphedema , Allergies
20798451	5RH 56.3(UtSMC)		
264487	5RH.I (Brain)	Brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
			disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy,
		ļ	Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
•			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264534	5RH.11 (Bone marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies, transplantation, Graft vesus host,
264535	5RH.12 (Bone marrow)	Bone Marrow	Hemophilia, hypercoagulation, idiopathic
204333	Jan. 12 (Dolle Hallow)		thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies, transplantation, Graft vesus host,
	,		
264563	5RH.19 (Fetal Brain)	Fetal brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
20.22	,		disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy,
		ŀ	Epilepsy, Lesch-Nyhan syndrome, Multiple
		1	sclerosis,Ataxia-
	}	-	telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264488	5RH.2 (Bone Marrow)	Вопе Магтоw	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies,
		ľ	immunodeficiencies, transplantation, Graft vesus host,
264564	5RH.20 (Lymph Node)	Lymph Node	Lymphedema , Allergies
264565	5RH.21 (Pancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
264566	5RH.22 (Placenta)	Placenta	Infertility, birth defects
264567	5RH.23 (Thyroid)	Thyroid	Hyperthyroidism and Hypothyroidism
264591	5RH.25 (Fetal Brain)	Fetal brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
204371	,		disease, Stroke, Tuberous sclerosis, hypercalceimia,
		r	Parkinson's disease. Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
	1 .		sclerosis,Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264592	SRH.26 (Bone Marrow)	Вопе Магтом	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies, transplantation, Graft vesus host,
264593	5RH.27(thyroid)	Thyroid	Hyperthyroidism and Hypothyroidism
264594	5RH.28 (Pancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
264595	5RH.29 (Lymph Node)	Lymph Node	Lymphedema , Allergies
264489	5RH.3 (Bone Marrow)	Bonc Marrow	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies, transplantation, Graft vesus host,
ł		1	

264596	5RH.30 (Placenta)	Placenta	Infertility, birth defects
264628	SRH.33 (fetal Kidney)	Fetal kidney	Diabetes, Autoimmune disease, Renal artery stenosis,
			Interstitial nephritis, Glomerulonephritis, Polycystic
	j		kidney disease, Systemic lupus erythematosus, Renal
	•		tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch-
			Nyhan syndrome
264629	5RH.34 (lymph Node)	Lymph Node	Lymphedema, Allergies
264630	5RH.35 (Pancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
264631	SRH.36 (thyroid)	Thyroid	Hyperthyroidism and Hypothyroidism
264632	5RH.37 (Fetal Brain)	Fetal Brain	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
			disease, Stroke. Tuberous sclerosis, hypercalceimia,
			Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
	<u> </u>		sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264490	5RH.4 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
			thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies, transplantation, Graft vesus host,
264681	SRH.43.1 (fetal thymus -	Fetal Thymus	Hemophilia, hypercoagulation, Idiopathic thrombocytopenic purpura, immunodeficiencies
	CRL7046)		Leukernia, osteoporosis, post-chemotherapeutic stern cell
264682	5RH.43.2 (hematopoetic stem	Hematopoeitic stem cells	repopulation
	celis - CRL2043)		
264683	5RH.43.3 (osteogenic sarcoma cell lines - HTB96)	Osteogenic Sarcoma	Sarcomas, osteoporosis, osteopetrosis
264684	5RH.43.4 (Fetal Liver)	Fetal Liver	Von Hippel-Lindau (VHL) syndrome,
			Cirrhosis, Transplantation
264685	5RH.43.6 (Spleen)	Spleen	Hemophilia, Hypercoagulation, Idiopathic
			thrombocytopenic purpura, Immunodeficiencies, Graft vesus host
			Von Hippel-Lindau (VHL) syndrome , Alzheimer's
264686	SRH.43.7 (pituitary)	Pituitary	disease, Stroke, Tuberous sclerosis, hypercalceimia,
	1		Parkinson's disease, Huntington's disease, Cerebral palsy,
	ļ		Epilepsy, Lesch-Nyhan syndrome, Multiple
	İ		sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
264757	SRH.44.1 (Kidney)	Kidney	Diabetes, Autoimmune disease, Renal artery stenosis, Interstitial nephritis, Glomerulonephritis, Polycystic
	1		kidney disease, Systemic lupus erythematosus, Renal
			tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch-
			Nyhan syndrome
		result inno	Von Hippel-Lindau (VHL) syndrome,
264759	5RH.44.2 (Fetal Liver)	Fetal Liver	Cirrhosis, Transplantation
264251	(6DU 44.2 (Usess)	Heart ·	Cardiomyopathy, Atherosclerosis, Hypertension,
264761	5RH.44.3 (Heart)	I see t	Congenital heart defects, Aortic stenosis, Atrial septal
			defect (ASD), Atrioventricular (A-V) canal defect, Ductus
			arteriosus, Pulmonary stenosis, Subaortic stenosis,
			Ventricular septal defect (VSD), valve diseases, Tuberous
<b> </b> -	İ		sclerosis, Scleroderma, Obesity, Transplantation
264763	5RH.44.4 (Prostate)	Prostate	Prostate Cancer
264765	5RH.44.5 (Spleen)	Spieen	Hemophilia, Hypercoagulation, Idiopathic
			thrombocytopenic purpura, Immunodeficiencies, Graft
i	I	i	- vesus host

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		(a)	Von Hippel-Lindau (VHL) syndrome , Alzheimer's
264767	5RH.44.6 (Pituitary)	Pituitary	disease, Stroke, Tuberous sclerosis, hypercalceimia,
			Parkinson's disease. Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
	İ		
		·	scierosis, Ataxia- telangiectasia, Leukodystrophies, Behavioral disorders,
	l		Addiction, Anxiety, Pain, Neuroprotection
264828	5RH.46.1 (Lymph Node)	Lymph Node	Lymphedema, Allergies
264887	5RH.47.5 (Fetal Liver)	Fetal Liver	Von Hippel-Lindau (VHL) syndrome,
			Cirrhosis, Transplantation
18108377	5RH.50.1 (B's lymphoma)	Burkitt's Lymphoma	Lymphoma, blood cancers
18108380	5RH.50.2 (thalamus)	Thalamus	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
			disease, Stroke, Tuberous sclerosis, hypercalceimia.
	1		Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis,Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
18108396	5RH.50.3 (adrenal gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia,
10100201	SDEL SO A (fetal lung)	Fetal Lung	Airway diseases, infection
18108391	SRH.50.4 (fetal lung) SRH.50.5 (salivary gland)	Salivary Gland	Dry mouth, infection
18108357	SRH.50.6 (mammary gland)	Mammary Gland	Lactation disorders, breast cancer
18108390		Bone Marrow	Hemophilia, hypercoagulation, Idiopathic
264532	5RH.9 (Bone Marrow)	Bone Mariow	thrombocytopenic purpura, autoimmume disease, allergies,
			immunodeficiencies, transplantation, Graft vesus host,
			immunodenteleites, manspiantation, Olan 1998 11229
263974	736xN		
263976	736xN		
263981	736xN		
20281166	96xN		
20281169	96xN		
20281171	96xN		
263994	cDNA-ORF Selection		
264080	Mx96		
21906754	NOH 6.1 (HH729)		
22278996	NOH 6.10 (PrEC)	Endothelial cells	heart disease, cancer
22278997	NOH 6.11 (CAEC)	Endothelial cells	heart disease, cancer
22278998	NOH 6.12 (CSC)	Cancer Cell line	Cancer
22278999	NOH 6.13 (NHNPC)	Cancer Cell line	Cancer
22279000	NOH 6.14 (NHMC-RM)	Cancer Cell line	Cancer
22279002	NOH 6.15 (Hypothalmus)	Hypothalamus	Von Hippel-Lindau (VHL) syndrome, Alzheimer's
	, , , , , , , , , , , , , , , , , , , ,		disease, Stroke, Tuberous sclerosis, hypercalceimia,
,			Parkinson's disease, Huntington's disease, Cerebral palsy,
	1	1	i antilibert a creating the contract of the co
			Epilepsy, Lesch-Nyhan syndrome, Multiple
,		*	Epilepsy, Lesch-Nyhan syndrome, Multiple
,		÷	Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxia-
21006761	NOU C 2 (In David Blatalana)	Distalare	Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxia-telangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection, Obesity
21906764	NQH 6.2 (In Dated Platelets)	Platelets	Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxia-telangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection, Obesity Clotting diseases, stroke
21906764	NQH 6.3 (HuVec)	Endothelial cells	Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxia- telangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection, Obesity  Clotting diseases, stroke heart disease, cancer
			Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxia- telangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection, Obesity Clotting diseases, stroke heart disease, cancer heart disease, cancer
21906765	NQH 6.3 (HuVec) NQH 6.3 (Sized-HUVEC) NQH 6.4 (UtMVEC- myo)	Endothelial cells Endothelial cells Cancer Cell line	Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxia- telangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection, Obesity Clotting diseases, stroke heart disease, cancer heart disease, cancer Cancer
21906765 87168474	NQH 6.3 (HuVec) NQH 6.3 (Sized-HUVEC)	Endothelial cells Endothelial cells Cancer Cell line Cancer Cell line	Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxia- telangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection, Obesity  Clotting diseases, stroke  heart disease, cancer heart disease, cancer Cancer Cancer
21906765 87168474 21906766	NQH 6.3 (HuVec) NQH 6.3 (Sized-HUVEC) NQH 6.4 (UtMVEC- myo) NQH 6.5 (NHEM-neo) NQH 6.6 (NHEK)	Endothelial cells Endothelial cells Cancer Cell line	Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxia- telangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection, Obesity  Clotting diseases, stroke  heart disease, cancer heart disease, cancer Cancer Cancer Cancer Cancer
21906765 87168474 21906766 21906767	NQH 6.3 (HuVec) NQH 6.3 (Sized-HUVEC) NQH 6.4 (UtMVEC- myo) NQH 6.5 (NHEM-neo)	Endothelial cells Endothelial cells Cancer Cell line Cancer Cell line	Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxia- telangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection, Obesity  Clotting diseases, stroke  heart disease, cancer heart disease, cancer Cancer Cancer

WO 00/58473

22278995	NQH 6.9 (PrSC)	Cancer Cell line	Cancer
27486261	NQH 7.1 (Jurkat E6-	Cancer Cell line	Cancer
	untreated)		
27486262	NOH 7.2 (TF1-untreated)	Cancer Cell line	Cancer
27486264	NQH 7.3 (U87-untreated)	Cancer Cell line	Cancer
27486265	NOH 7.4 (THP1-untreated)	Cancer Cell line	Cancer
29331822	NQH 8.1 (Brain- amygdala)		Von Hippel-Lindau (VHL) syndrome, Alzheimer's
2,55.022	The state of the s		disease, Stroke, Tuberous sclerosis, hypercalceimia,
	•		Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
	1		telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
29331824	NQH 8.2 (Brain-		Von Hippel-Lindau (VHL) syndrome, Alzheimer's
29331024	hippocampus)		disease, Stroke, Tuberous sclerosis, hypercalceimia,
	inppocaripus,		Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
			sclerosis, Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
			Addiction, Anxiety, Pain, Neuroprotection
20221026	NOH 8.3 (Brain- substantia		Von Hippel-Lindau (VHL) syndrome, Alzheimer's
29331825	• '		disease, Stroke, Tuberous sclerosis, hypercalceimia,
	nigra)		Parkinson's disease, Huntington's disease, Cerebral palsy,
			Epilepsy, Lesch-Nyhan syndrome, Multiple
		:	sclerosis.Ataxia-
			telangiectasia, Leukodystrophies, Behavioral disorders,
	1		Addiction, Anxiety, Pain, Neuroprotection
		0 11:	digestive diseases, obesity, diabetes
29331826	NQH 8.4 (small intestine)	Small intestine Spinal chord	paralysis, neurodegenerative disorders
29331827	NQH 8.5 (Spinal cord)	Stomach	Stomach cancer
29331828	NQH 8.6 (stomach)	Trachea	Airway diseases, infection
29331830	NQH 8.7 (Trachea) NOH 9.1 (Sized-MG-	Haciica	
87168518	63 treatment pool)		
87168559	NOH 9.2 (Sized-HEPG2		
8/108339	untreated)		
35695763		Cancer Cell line	Cancer
33093703	Training (more remains)		
35695855	NQH.10.2 (U-937_treatment	Cancer Cell line	Cancer
33093833	pool)		
35695917	NQH.10.3 (JAR)	Cancer Cell line	Cancer
35696052	NOH.10.4 (PA-1)	Cancer Cell line	Cancer
35696286	NOH.10.5 (CADMEC)	Endothelial cells	heart disease, cancer
35696423	NQH.10.6 (CADMEC_LA)	Endothelial cells	heart disease, cancer
52644045	NQH.11.1 (SK-PN-DW)	Cancer Cell line	Cancer
52644150	NQH.11.2 (Chorionic Villus	Chorionic villus	fertility, birth defects
22077130	Cells)		
52644229	NQH.11.3 (A549)	Cancer Cell line	Cancer
52644296	NQH.11.4 (U266B1)	Cancer Cell line	Cancer
52644332	NQH.11.5 (Daoy)	Cancer Cell line	Cancer
52644507	NOH.11.6 (SW1783)	Cancer Cell line	Cancer
52645080	NQH.12.1 (U-118MG)	Cancer Cell line	Cancer
52645129	NOH.12.2 (A204)	Cancer Cell line	Cancer
52645156	NOH.12.3 (T24)	Cancer Cell line	Cancer
52646317	NQH.12.4 (G-401)	Cancer Cell line	Cancer
	NOH.12.5 (CaSki)	Cancer Cell line	Cancer
52646365 52646842	NOH.12.6 (SHP-77)	Cancer Cell line	Cancer
1 1 / DANEA /	HAGETTER (OTH - 11)	COLUMN CONTINUE	<u> </u>

60424179	NQH.14.1 (Yale75_breast carcinoma)	Breast carcinoma	Breast Cancer
60424269	NQH.14.2 (Yale78B_ovarytumor)	Ovary tumor	Ovarian cancer
60431528	NQH.14.3 (Yale79_prostateBPH)	Prostate	Prostate Cancer
60431602	NQH.14.4 (Yale80_ProstateAdenocarcin oma)	Prostate	Prostate Cancer
60431735	NQH.14.5 (Yale86_UterineMyoma)	Uterine Myoma	Uterine Cancer
60431850	NQH.14.6 (Yale207_Myometrium)	Myometrium	Fertility
60432049	NOH.15.1 (Yale99 cervix)	Cervix	Osteoporosis, cervical cancer
60432113	NQH.15.2 (Yale45_spleenITP)		Hemophilia, Hypercoagulation, Idiopathic thrombocytopenic purpura, Immunodeficiencies, Graft vesus host
60432229	NQH.15.3 (Yale16_Skin)	Skin	wound healing, melanoma
60432289	NQH.15.4 (Yale137_Parotid)		·
60433356	NQH.15.5 (Yale38 SmallIntestine)	Small intestine	digestive diseases, obesity, diabetes
60433438	NQH.15.6 (Yale28_ColonAscending)	Colon	Colon cancer
65274444	NOH.17.1 (Larynx)	Larynx	Cancer
65274572	NQH.17.2 (Duodenum)	Duodenum	
65274620	NQH.17.3 (Kidney, Primary tumors)		Diabetes, Autoimmune disease, Renal artery stenosis, Interstitial nephritis, Glomerulonephritis, Polycystic kidney disease, Systemic lupus erythematosus, Renal tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch- Nyhan syndrome
65274727	NQH.17.4 (Lung Pleura, normal)	Lung	Airway diseases, infection
65274791	NQH.17.5 (Lung, Normal Adult)	Lung	Airway diseases, infection
83373044	NQH.18.230 (Pooled adrenal gland, placents)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia
85658542	NQH.18.560 (Pooled uterus, BeWo pool)	Uterus	Infertility, birth defects
33656970	NQH.9.1 (MG-63_treatment pool)	Cancer Cell line	Cancer
33657023	NQH.9.2 (HEPG2 untreated)		Von Hippel-Lindau (VHL) syndrome, Cirrhosis,Transplantation
33657084	NQH.9.3 (PC3_untreated)	Cancer Cell line	Cancer
33657109	NQH.9.4 (TF-1_TPA)	Cancer Cell line	Cancer
33657182	NQH.9.5 (TF-1_TPO)	Cancer Cell line	Cancer
33657349	NQH.9.6 (TF-1_Hernin)	Cancer Cell line	Cancer
33657402	NQH.9.7 (HFDPC)	Cancer Cell line	Cancer
264259	NQHI (Mixture of eight adult & two fetal tissues)		
264288	NQH2 (Ten tissues plus lymphocyte control)		
264448	NQH3 (Bone Marrow)	Bone Marrow	Hemophilia, hypercoagulation, idiopathic thrombocytopenic purpura, autoimmume disease, allergies, immunodeficiencies, transplantation, Graft vesus host,
265017	NQH4.1 (lymph node)	Lymph Node	Lymphedema, Allergies

265018	NQH4.2 (fetal kidney)	Fetal Kidney	Diabetes, Autoimmune disease, Renal artery stenosis, Interstitial nephritis, Glomerulonephritis, Polycystic kidney disease, Systemic lupus erythematosus, Renal tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch-Nyhan syndrome
66712502	NQH4.2 (Sized)		
265019	NQH4.3 (pituitary gland)		Von Hippel-Lindau (VHL) syndrome, Alzheimer's disease, Stroke, Tuberous sclerosis, hypercalceimia, Parkinson's disease, Huntington's disease, Cerebral palsy, Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxiatelangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection, Obesity
66714117	NQH4.3 (Sized)		
265020	NQH4.4 (testis)	testis	Infertility, birth defects
265021	NQH4.5 (fetal liver)	Fetal Liver	Von Hippel-Lindau (VHL) syndrome, Cirrhosis, Transplantation
265022	NQH4.6 (thyroid)	Thyroid	Hyperthyroidism and Hypothyroidism
18108376	NQHS.1 (MCF-7)	Breast cancer	Breast Cancer
18108387	NQH5.2 (CCRF-CEM)	Cancer Cell line	Cancer
264952	NRL1: HPLC FRACTIONATION OF RE- LIG		
263971	Old BB3 Baits		
263969	Old BB5 Baits		
263975	ORFSEL		
263972	OTHER Baits		
263978	pGALORF		
264106	PPBAITS		
264088	QC-YA7		
264089	QC-YA8		
264102	Resequenced Interactors	•	
264369	RRH.1		
60170394	RRH.10.1 (MCF-7untreated)	Breast cancer	Breast Cancer
60170615	RRH.10.2 (U-937_treatment pool)	Cancer Cell line	Cancer
60170831	RRH.10.3 (JAR)	Cancer Cell line	Cancer
60174639	RRH.11.8 (HeLa)	Cancer Cell line	Cancer
264113	πQEA Baits		
263973	RRQEA_B5 baits		
29146498	SRD 3.1 (SKMC)	Cancer Cell line	Cancer
29146499	SRD 3.2 (SKMC)	Cancer Cell line	Cancer
29147620	SRD 3.3 (RPTEC)	Cancer Cell line	Cancer
29148627	SRD 3.4 (HRCE)	Cancer Cell line	Cancer
29148629	SRD 3.6 (HRE)	Cancer Cell line	' Cancer
29148784	SRD 3.7 (HRE)	Cancer Cell line	Cancer
55810764	SRD.7.1 (Lymph Node)	Lymph Node	Lymphedema, Allergies
55811150	SRD.7.2 (pancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
55811386	SRD.7.3 (Adrenal Gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia,

55811576	SRD.7.4 (Pituitary Gland)	Pituitary	Von Hippel-Lindau (VHL) syndrome, Alzheimer's disease, Stroke, Tuberous sclerosis, hypercalceimia, Parkinson's disease, Huntington's disease, Cerebral palsy, Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxiatelangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection, Obesity
55811957	SRD.7.5 (Fetal Liver)	Fetal Liver	Von Hippel-Lindau (VHL) syndrome, Cirrhosis, Transplantation
55812038	SRD.7.6 (Fetal Kidney)	Fetal kidney	Diabetes, Autoimmune disease, Renal artery stenosis, Interstitial nephritis, Glomerulonephritis, Polycystic kidney disease, Systemic lupus erythematosus, Renal tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch- Nyhan syndrome
56181562	SRD.8.1 (Lymph Node)	Lymph Node	Lymphedema, Allergies
56181686	SRD.8.2 (Pancreas)	Pancreas	Pancreatitis, diabetes, pancreatic cancer
56182181	SRD.8.3 (Adrenal Gland)	Adrenal Gland/Suprarenal gland	Adrenoleukodystrophy , Congenital Adrenal Hyperplasia,
56182323	SRD.8.4 (Pituitary Gland)	Pituitary	Von Hippel-Lindau (VHL) syndrome, Alzheimer's disease, Stroke, Tuberous sclerosis, hypercalceimia, Parkinson's disease, Huntington's disease, Cerebral palsy, Epilepsy, Lesch-Nyhan syndrome, Multiple sclerosis, Ataxiatelangiectasia, Leukodystrophies, Behavioral disorders, Addiction, Anxiety, Pain, Neuroprotection, Obesity
56182435	SRD.8.5 (Fetal Liver)	Fetal Liver	Von Hippel-Lindau (VHL) syndrome, Cirrhosis, Transplantation
56182575	SRD.8.6 (Fetal Kidney)		Diabetes, Autoimmune disease, Renal artery stenosis, Interstitial nephritis, Glomerulonephritis, Polycystic kidney disease, Systemic lupus erythematosus, Renal tubular acidosis, IgA nephropathy, Hypercalceimia, Lesch-Nyhan syndrome
32833986	SRD4: HL adapter		
56526486	SRD5.1:rr fragments		
33109954	SRD5: long-RXRJ		
56994075	SRD9.1 (CS/SC)	Cancer Cell line	Cancer
263977	TSC Screen 1	<u> </u>	

D34

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